

TreeAge Pro 2015 User's Manual

TreeAge Pro 2015 User's Manual

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1. Introduction

This chapter serves as an introduction to TreeAge Pro and TreeAge Software, Inc.

1.1 Welcome to TreeAge Pro

TreeAge Pro has been designed to implement the techniques of decision analysis in an intuitive and easy-to-use manner. It transforms decision analysis from a potentially tedious exercise into an easily applied and highly visual means of:

1. *Organizing* the decision making process;
2. *Analyzing* the problem at hand; and
3. *Communicating* the structure of the problem, the nature of the uncertainties, and the basis for a strategy recommendation.

If you are experienced in decision analysis, you will find TreeAge Pro easy to use following only a cursory review of the software commands, although the richness of the program will become more apparent with further study of the manual.

If you are looking for features added to recent releases, please refer to the What's New in TreeAge Pro section of this chapter.

If you have no, or only limited, experience with decision analysis, TreeAge Pro will make it much easier to learn. Go to the Decision Analysis Primer chapter for a basic decision analysis primer.

The organization of the rest of the manual is described in the last section of this Introduction Chapter.

1.2 TreeAge Pro Interface

TreeAge Software embarked on an ambitious project to rewrite the software within the Eclipse development environment. Eclipse is a customizable development environment that allowed us to incorporate many of its native features and third-party plugins into TreeAge Pro.

This new architecture not only improves the user interface and other components of TreeAge Pro 2011 and subsequent versions, but it also better positions us to add more functionality in future releases.

We hope you find that the newest version of TreeAge Pro is the most flexible, user-friendly version of the software released to date. Prior users of TreeAge Pro will notice a big change in the TreeAge Pro Interface.

For an introduction of the new interface, please refer to the TreeAge Pro Interface chapter.

1.3 Optional modules

TreeAge Pro follows a modular design. The "Core" module includes both decision tree functionality, Monte Carlo simulation, multi-way sensitivity analysis, and much more. Additional features may be added in the form of two optional add-ons: a Healthcare module and an Excel™ add-in module.

The TreeAge Pro Healthcare module integrates core decision analysis functionality with specialized capabilities, including:

- Markov cohort analysis
- Individual-level, Markov simulation
- Cost-Effectiveness analysis
- Net Benefits analysis
- Additional information available at <https://www.treeage.com/shop/treeage-pro-healthcare/>

Healthcare Module Functions

The TreeAge Pro Excel module can help:

- Automate the creation of Excel charts and reports from within TreeAge Pro
- Simplify the integration of spreadsheets and TreeAge Pro models
- Edit input data in Excel
- Output analysis data to Excel
- Additional information available at <https://www.treeage.com/shop/treeage-pro-excel/>

Excel Module Functions

The TreeAge Pro Suite includes all modules, including Healthcare and Excel.

1.4 Integration with other software

In TreeAge Pro, users of the Excel module gain access to powerful features that allow automation of TreeAge Pro tasks via scripts and macros, in order to control the updating and analysis of your models.

In addition, the embedded Python script interpreter has been improved, giving modelers easy access to the powerful Python programming language.

1.5 Installation and system requirements

To install TreeAge Pro, please go to www.treeage.com/install and enter your serial number. You will be presented with a list of installers. Choose the one that is appropriate for your computer. After installing, you will need to authorize TreeAge Pro with your license serial number.

System requirements are available at <https://www.treeage.com/support/system-requirements/>.

1.6 Cross-Platform Compatibility

TreeAge Pro 2011 and subsequent versions are built in the Eclipse architecture, which allows the software to run on Windows, Mac, and Linux operating systems. There are different installers for each OS available at our website. In each OS, the software should have a look and feel that for the most part matches the native view for that OS.

Since TreeAge Pro 2009 and earlier versions ran only on the Windows OS, most of the instructions in this manual describe Windows operations. There are some differences in commands/mouse-clicks on Mac and Linux. The table below describes some of these differences.

Windows	Mac	Linux
Control + key	Command + key	?
Right-click	Control-click	
Function keys	Fn key + function key	

Command differences among operating systems

1.6.1 Mac Function Keys

Mac computers often use function keys to interact with the Mac Operating System. You can release those function keys from their OS functions by changing settings in the System Preferences under the category Keyboard. Specifically...

- Check the box to use F1, F2, etc. as standard function keys.
- Uncheck the box for active use of "Expose & Spaces" or "Mission Control".

1.6.2 Linux with GNOME Desktop Environment

If you use Unix-like operating systems with the GNOME desktop environment, you may be unable to see the application's menu icons.

If you experience this problem:

- Start the gconf-editor.
- Select the checkbox in /desktop/gnome/interface/menus_have_icons

1.7 Get technical support

There are several ways to get help in using TreeAge Pro.

1.7.1 Using the online help

TreeAge Pro installs with the online help you are currently reading. This help is quite extensive and describes both modeling techniques and technical aspects of TreeAge Pro.

1.7.2 Web site resources

The support section of our web site includes a variety of information for users of the software:

<http://www.treeage.com/support>

There are also product demos available:

<http://www.treeage.com/training>

1.7.3 Example models

Most chapters include tutorials and references to example models, which are installed on your computer with TreeAge Pro. The Managing Projects and Documents Chapter describes how to access these models.

1.7.4 E-mail and telephone support

If you are a registered user of TreeAge Pro with a current maintenance agreement, you are eligible for free e-mail and telephone support; limited technical support is available if you are evaluating the trial software. We would like to make your use or evaluation of the software as productive and pleasant as possible. If you need help:

- Send an e-mail to support@treeage.com.
- Call us at 1-413-458-0104, then press 2 for Support.
(1-888-TREEAGE in US/Canada)

Access to technical support

1.8 File compatibility

TreeAge Pro 2011 introduced a new XML-based model file structure.

- Models saved in TreeAge Pro 2011 or later use the file extension .trex.
- Models saved in TreeAge Pro 2009 or earlier use the file extension .tre or .pkg.

File formats

1.8.1 Compatibility issues with new features

New features have been introduced which are not supported by older versions of the post-2011 software. Models that do not use those features can be opened in the older post-2011 software without issue.

Model validation allows you to identify features that will not work in older post-2011 software. Use of those features must be removed if sharing with a colleague with older software.

1.8.2 Compatibility between models from TP 2009 and TP 2011

Older *.tre and *.pkg files can be opened in TreeAge Pro 2011 and subsequent versions. When you open a files from an older format, you are asked whether you want to save the file in the new format or the old format. Not all features of TreeAge Pro 201x can be saved in the TreeAge Pro 200x format.



It is recommended that the old *.tre files are first saved as *.pkg files before opening them in TreeAge Pro 201x. That way, the tables used by the model will be included with the new *.trex document.

TreeAge Pro 201x can open trees created in previous versions of TreeAge Pro, as well as in DATA Pro or DATA. However, unless your TreeAge Pro software includes the Healthcare module, access to Markov and cost-effectiveness models created in either DATA or TreeAge Pro will be limited to viewing the file.

Trees created in TreeAge Pro 201x are not directly backward-compatible with TreeAge Pro versions. However, you can save models in an older format.

To save a model in an older TreeAge Pro format:

- Choose File > Save As from the menu.
- Select the file type TreeAge Pro 200x Package (*.pkg).
- Save the file.

Certain features not present in the earlier software will be lost in the conversion and, in any event, care should be taken following conversion to confirm the accuracy of calculations.

1.9 Manual structure

This Manual is broken into several groups of chapters.

1. Getting Started with TreeAge Pro (starting with this chapter)
2. Working with Decision Trees (starting with the Analyzing Decision Trees Chapter)
3. Uncertainty and Variability: Using Variables, Tables, and Distributions (starting with the Introduction to Variables and Sensitivity Analysis Chapter)

4. More Decision Tree Modeling and Analysis Options (starting with the Stored Analysis Abstracts and Sequences Chapter)
5. Working with the Excel/COM Module (starting with the Connecting Spreadsheets and Trees Using the Excel Module Chapter)
6. Cost-Effectiveness Analysis with Tree Age Pro Healthcare & TreeAge Pro Suite (starting with the Building and Analyzing Cost-Effectiveness Models Chapter)
7. Markov Modeling, Analysis, and Simulation with the Healthcare Module (starting with the Building and Analyzing Markov Models Chapter)

Sections of this manual

1.10 What's New in TreeAge Pro

This section identifies new features in the latest releases.

1.10.1 TreeAge Pro 2015, Release 1

The following features were included in TreeAge Pro 2015, Release1:

- Monte Carlo simulation performance has been significantly improved through the incorporation of a faster database engine and more efficient data storage/queries and by allowing simulation data to be stored on disk rather than exclusively in memory. Not only will your simulations run faster, you will also be able to run larger simulations than previously possible.
- New perspectives/layouts (Build, Analyze) arrange the application window for efficient work and provide toolbar icons for commonly used functions.
- TreeAge Pro now prompts you for data input of node labels, branch probabilities, payoff entries, etc. to simplify data entry within the model. This is highlighted in the Decision Tree Tutorial Chapter.
- TreeAge Pro's formula editor has been improved to make it easier to build numeric expressions within the model.
- The Configure Model wizard walks you through the key Tree Preferences categories to quickly configure your model for proper calculation and display.
- The Model Dashboard presents an "at a glance" view of model configuration and model inputs with immediate access to edit these elements.
- Global Discounting allows you to discount all entires for each payoff/reward set with a single change to the Tree Preferences.
- Create a Bilinks Report to review the connections between your model and an Excel workbook.
- Better identify optimal, rejected and excluded strategies when a model is rolled back.
- Multi-attribute weights can now be entered using variables and expressions.
- Hide/show all notes in the tree editor via Tree Preferences.
- Restructured example model folders to make it easier to find example models in the Projects View.

- More efficiently export large simulation data sets through a new more efficient export option.
- Quickly copy graph images for pasting into documents.

1.10.2 TreeAge Pro 2014, Release 2

The following features were included in TreeAge Pro 2014, Release 2:

- Sensitivity analysis now supports adjustments up and down relative to the parameter base value.
- Cost-effectiveness sensitivity analysis output now includes net monetary benefit for each strategy. This allows you to more easily identify thresholds in the main output.
- New functions support continuous time discounting within discrete event simulation (DES) models.
- The Add/Edit Distribution dialog now provides access to the Help files for access to distribution formulas, which may be helpful when entering distribution parameters.
- Filtered simulation output now shows the count of filtered rows relative to the entire dataset.
- New global matrix functions differentiate between data storage, retrieval, increment, initialize and export. The separate functions allow better Help for proper function syntax.
- Models now support custom functions for specific commands/formulas. Pass different arguments to the functions to use the custom function in different contexts.
- New back/forward toolbar buttons allow you to move among the open tabs for models and analysis output.
- Perspectives/layouts were updated to better group related views in the same pane within the application window. To use the updated layouts, you must create a new workspace.
- The bottom of the application window now displays node counts, both for the whole model and for the selected node and its subtree.
- A new tree preference option allows you to show Markov and DES rewards for all enabled reward sets rather than just the reward sets selected for the calculation method.
- New model validation rule checks for consistent use of half-cycle correction.
- New keyword `_node_id` provides access to the identifier for the current node. This data can be stored in global matrices for patient tracking.

1.10.3 TreeAge Pro 2014, Release 1

The following features were included in TreeAge Pro 2014, Release 1:

- Time-to-Event simulation: New node types and functions were added to facilitate time-to-event simulation.
- Monte Carlo output filtering: Filter output data based on user-defined criteria. Change active payoffs without rerunning the analysis. Generate secondary reports based on the filtered data.
- Performance: Enhanced software to improve viewing and editing performance when working with large models.

- Clones: View the full read-only structure of clone copies. When errors occur within the expanded clone copy, highlight the proper node in the clone copy. This option had been disabled in prior software versions.
- Analysis keywords: Added/improved analysis keywords. Some keywords allow execution of commands after analyses or before/after specific iterations within simulations. Other keywords accumulate rewards within the analysis within time-to-event simulations to drive model flow or verify/debug models.
- Monte Carlo output reports: Hide PSA-specific secondary that are only valid for Probabilistic Sensitivity Analysis from Microsimulation results.
- Enhanced branch and path probability functions: BranchProb and PathProb functions can now be used within the model rather than only at terminal nodes.
- Restructured categories in model input views: Improved UI to use less space for categories except when specifically working with categories.

1.10.4 TreeAge Pro 2013, Release 2

The following features were included in TreeAge Pro 2013, Release 2:

- Simple interface option: A simple interface "skin" helps new users focus on tree modeling and analysis basics, speeding up the learning process. A new Welcome View provides new users with support, training and demo resources directly from the initial presentation window.
- Cost-effectiveness analysis: CEA graphs and reports have been enhanced to better highlight the favored strategy.
- Function Helper: Use the Function Helper to see microhelp for built-in functions and as a guide for function argument entries.
- Node-level Probability Coherence Override: Allow non-coherent probabilities or normalize probabilities at specific chance and Markov nodes.
- Summation Nodes: Use a summation node when the values of a nodes branches need to be combined together to calculate the total value of the summation node.
- Variable Definition Entry: Enter complex variable definitions in the Add/Change Variable dialog.
- BilinkName Function: Reference Bilinks by name rather than by index.
- File Compatibility: Use model validation to identify features that require new or recent versions of TreeAge Pro before sending a model to a colleague.

1.10.5 TreeAge Pro 2013, Release 1

The following features were included in TreeAge Pro 2013, Release 1:

- State Transition Diagrams: Create a state transition diagram to represent a Markov model. Convert the state transition diagram to the decision tree format. Convert your completed Markov model in decision tree format back to a state transition diagram.

- Influence Diagrams: Create an influence diagram as a combination of decision, chance, deterministic and value nodes. The nodes are connected by arcs which represent the time relationship among the events represented by the nodes. Convert the influence diagram to a decision tree.
- Strategy Exclusion: Exclude specific strategies from all analyses.
- Model Validation: Identify and highlight common model flaws.

1.10.6 TreeAge Pro 2012, Release 2

The following features were included in TreeAge Pro 2012, Release 2.0:

- Bayes Revision: Revise probabilities using Bayes' Theorem when your model includes imperfect tests. This feature handles both sensitivity/specificity and a general m by n grid.
- Enhanced Search/Replace with Highlighting: This enhancement allows you to view and/or replace the search results in a structured list. Search results are also highlighted in the model structure.

The following features were included in Release 2.1:

- Object Interface Enhancements: The programming interface has been modified and enhanced to provide better access to modeling objects and analysis output. It is now possible to access secondary analysis output from Monte Carlo simulations and sensitivity analyses.
- Highlighting Expressions: The Variable Properties, Tracker Properties, Distributions and Tables Views were enhanced to allow you to highlight the use of a model input within the Tree Diagram Editor.
- Image Exports: Generate higher quality images of graphs and models for presentation and publication.
- Improved User Interface Presentation: the Tree perspective has been rearranged to show the most commonly used Views and to place similar views in the same area of the user interface. Right-click on the Tree perspective and choose Reset to use the new format.

1.10.7 TreeAge Pro 2012, Release 1

This release introduced the following features:

- Sensitivity Analysis and Microsimulation: Run one-way sensitivity analysis and Microsimulation at the same time. Microsimulation mean values are used as the expected values for each variable value.
- Distributed Simulations: Use slave computers to speed up long simulations.
- Variable Definition Arrays: Create variable definitions with multiple expressions via an array. The individual expressions are then referenced by index.
- Exportable summary report output: Output provided by the Markov Cohort Summary Report and the Cost-Effectiveness Text Report are now more easily exported to Excel.

- Model Documentation: Create a Help HTML file to describe your model to colleagues.
- News Reader: Keep informed about product releases/updates, training, etc. via the TreeAge Pro News Reader.

TreeAge Pro 2012, R1.0 Feature List

1.10.8 TreeAge Pro 2011, Release 2

This release introduced the following features:

- The TreeAge Pro Player: Create a Model Interface to allow non-licensed users of TreeAge Pro to review and/or analyze your model within the TreeAge Pro Player.
- Cohort size option multiplied by rewards in Full Markov Cohort output.

TreeAge Pro 2011, R2.0 Feature List

1.10.9 TreeAge Pro 2011, Release 1

This was the first release of TreeAge Pro using Eclipse RCP. All functions were "new" in this release.

2. Decision Analysis Primer

This chapter introduces the core concepts and methods of decision analysis as implemented in TreeAge Pro 201x.

2.1 What is decision analysis?

Decision analysis — as it is taught and practiced today in diverse fields such as oil and gas exploration, business, healthcare, law, and engineering — is a systematic approach to decision making under uncertainty. The process is designed to help decision makers think clearly about the many elements of complex decisions, such as:

- the range of possible consequences of actions (or inaction)
- preferences among different sets of consequences
- the impact of complex, unpredictable systems and processes (e.g., markets, geological structure, health)
- the actions of others (e.g., consumers, competitors, regulators, patients)

The concepts and methods of decision analysis are uniquely suited to incorporating into the decision making process both what is known about a problem, and also what is uncertain. TreeAge Pro is used in two of the key steps in decision analysis, modeling and analysis.

2.1.1 Modeling

Using decision analysis, a complex problem can be disaggregated into smaller problems and elements, which can be easier to understand. These components are then employed in building a model of the problem's essential elements.

First, a set of alternatives is compiled. Then, events and other factors that may affect the outcome of an alternative are identified. A factor whose impact on the final outcome is not known at the time of the decision is referred to as an uncertainty. It can be represented as either a structural element of the model, or a parameterized distribution used in a formula in the model. Based on the decision maker's objectives, one or more attributes are selected to quantify preferences for the range of possible final outcomes (and, ultimately, to rank alternatives). For example, a monetary scale would be used to measure different project cost scenarios and then to rank alternatives.

Compared to basic spreadsheet analysis or statistical modeling, a particular strength of decision analysis is the intuitive, visual form of the model. The model may be either a decision tree or a dependency diagram, which are different means of visually representing the same problem.

A decision tree is a branching structure in which various node symbols are used to represent different kinds of events, including decisions and uncertainties. A node's branches represent the outcomes or

alternatives associated with that event. Every series of actions and outcomes is clearly represented with a distinct path.

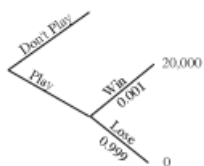
Alternatively, the same problem often may be represented more compactly by a dependency diagram. In a dependency diagram, each factor that directly or indirectly affects the final outcome is represented with a single node. Arcs between nodes show that one factor either affects another or precedes it in time.

2.1.2 Expected values and optimal decisions

While a model's explicit identification of the sequence and linkage of events is of great value in decision making, decision analysis is designed to do much more. Using basic concepts from probability theory and statistics, the decision maker can calculate a mean (or expected) value for each course of action. By calculating the value of each possible chain of events, and weighing uncertain results by the probability of each outcome, the decision maker can identify the sequence of decisions that will maximize value, minimize costs, or balance multiple attributes.

These calculations are commonly illustrated with a decision about playing a simple lottery, which can be thought of as a random variable having a simple probability distribution of outcomes as follows.

A \$20,000 lottery offers a 1-in-1000 chance of winning. The expected value of playing this lottery — ignoring the ticket price — equals $0.001 \cdot (20,000) + 0.999 \cdot (0)$, or \$20. Based on expected value, a reasonable decision would be to purchase a ticket if it costs \$20 or less.



Typically, decision analysis problems involve multiple uncertainties, with the outcome of the first lottery being a second lottery, and so on. The analysis of compound lotteries works backwards, calculating an expected value for the final lottery and using this result in the calculation of the prior lottery. When all lotteries have been resolved in this way, decisions are evaluated by optimization — picking the alternative with the best expected value.

2.1.3 Expected utilities and certainty equivalents

A decision maker's attitude towards *risk* can be incorporated into model calculations. Using the techniques outlined in utility theory, a choice between different lotteries can be made based on their expected utilities or certainty equivalents. These in turn depend on the decision maker's attitude towards risk, quantified using a utility function.

2.1.4 Sensitivity analysis and Monte Carlo simulation

As illustrated with the lottery example, one way in which decision analysis deals with uncertainty is to reflect it explicitly in the model's structure. Events which have a significant impact on outcomes, and which are not under the decision maker's control, can be described using chance nodes and incorporated into the model calculation. A problem may involve numerous uncertainties; not all of them can or should be represented in the structure of the model. To deal with this, deterministic sensitivity analysis and Monte Carlo simulation are used to examine the potential impact of parameter assumptions and other uncertainties.

Deterministic sensitivity analysis can take a variety of forms, including 1-, 2-, and 3-way sensitivity analysis and tornado diagrams; it can be used to identify critical uncertainties by examining the extent to which a model's calculations and recommendations are affected as a consequence of changing selected assumptions. Monte Carlo simulation, also referred to as probabilistic sensitivity analysis, can incorporate *all* parameter uncertainties. Probabilistic sensitivity analysis provides similar insights to deterministic sensitivity analysis, and can also quantify the level of confidence that can be placed in the model's results.

Sensitivity analysis, Monte Carlo simulation, and other analytical tools can also be used to improve decision making. This is done by determining the potential value of obtaining various kinds of information (*perfect, imperfect, or sampling information*) that might help resolve critical uncertainties.

2.2 A simple problem: How should I invest \$1000?

Now consider a simple example. You have \$1,000 to invest, and two potential investments: a volatile equity investment, and a risk-free certificate of deposit (CD). You will reconsider your investment decision at the end of one year, but not earlier. You have no fees or taxes to consider, just an identical \$1,000 investment in either case.

The CD pays simple interest at a rate of 5% annually — your return would be \$50.

In researching the equity investment, you have come up with a simple probability distribution describing its year-end performance: 1) a 30% probability that its market value will have gone up by \$500; 2) a 40% probability of a modest \$100 increase in value; and 3) a 30% probability of a substantial drop in value, -\$600. Your investment objective is to maximize growth, and you are sufficiently wealthy that the possible loss of \$600 does not pose a material threat.



Assigning 30%/40%/30% probabilities to the outcomes of the risky investment in the example follows a standard method for representing a probability distribution of outcomes based on expert opinion. This particular type of discrete distribution is referred to as a Swanson's mean, or 10/50/90, distribution. The three outcomes represent the 10th, 50th, and 90th percentile values elicited from the expert. A similar approach uses 25%/50%/25% probabilities for the three outcomes.

For a primer on applying the basic techniques of decision analysis to this problem, complete the rest of this chapter. If you are already familiar with the fundamentals of decision analysis, you may prefer to go directly to the Decision Tree Tutorial Chapter to learn how to build this model in TreeAge Pro 201x as a decision tree.

2.3 Decision trees

In building a decision tree, there are some basic guidelines to be considered:

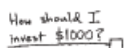
- *In the tree, events are ordered from left to right.* The tree often follows a time ordering of events, as outcomes become known to the decision maker. Time ordering is only critical, however, when a decision is made prior to knowing the outcome of a different event or when the probabilities of one event are dependent on another.
- *Different kinds of events are distinguished using shapes called “nodes.”* A decision node (square) indicates a choice facing the decision maker. A chance node (circle) represents an event which has multiple possible outcomes and is not under the decision maker’s control. A terminal node (triangle) denotes the endpoint of a scenario.
- *Branches “sprouting” from a decision node represent the set of actions being considered.* Decision alternatives are not required to be mutually exclusive (for example, one set of options could be “install smoke detectors,” “install fire extinguishers,” and “install smoke detectors and fire extinguishers”).
- *Branches from a chance node represent the set of possible outcomes of the event.* The branches must be mutually exclusive and exhaustive — in other words, defined such that all possibilities are covered and none overlap. Their probabilities must sum to 1.0 (100%).
- *Terminal nodes are assigned a value, referred to generically as a payoff.* All right-most nodes (those without branches) must be terminal nodes and have a payoff representing the net value — e.g., profit, cost, or utility — of that particular scenario’s series of actions and events.

Using these guidelines, let’s design a decision tree that represents the investment problem posed above.

2.3.1 Tree structure

The first event, the decision between the available investment options, is represented by a square (decision) node. This is the *root node* of the tree. It is labeled using a branch line to its left.

How should I
invest \$1000?

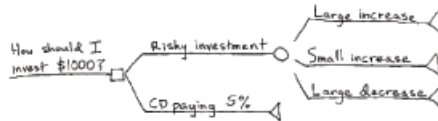


The two branches “sprouted” from the right side of the decision node represent the alternatives under consideration: (1) *Risky investment* and (2) *CD paying 5%*. Since there is no risk associated with the CD, this action really represents a final outcome, and a terminal node is drawn at the end of its

branch. For the risky equity investment, however, we decided that there were three possible outcomes, therefore Risky investment is drawn as a chance node.



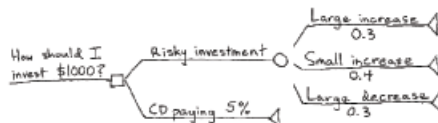
The three outcomes of the risky investment, Large increase, Small increase, and Large decrease, are drawn as branches emanating from the chance node. They are final outcomes, and are represented using terminal nodes.



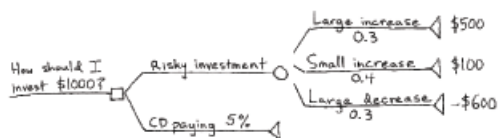
2.3.2 Probabilities and payoffs

Now the structure of the tree is complete. All that remains is to place the probabilities and payoff values in the tree.

Probabilities must be assigned to the branches emanating from a chance node, with the branch probabilities summing to 1.0. The probability for a particular outcome is indicated below the appropriate branch line.



Payoffs are assigned at every terminal node, and appear to the right of the node.



2.3.3 Calculating the tree

The investment decision tree is now complete, and can be evaluated. Decision tree calculations work backward, from right to left. Thus, calculating a tree is often referred to as “folding back” or “rolling back” the tree. The value of each node is determined as follows:

- The value of a terminal node is its assigned payoff value.
- The value of a chance node is its expected value. As described earlier, this is the mean value of the probability distribution specified by the chance node's branches.
- The value of a decision node is equal to its best option.

Now let's apply these rules to the investment tree.

The payoffs of the three terminal nodes are used. Working backward, starting from the topmost terminal nodes, you can find the expected value (EV) of the Risky investment chance node as follows:

$$= (500 * 0.3) + (100 * 0.4) + (-600 * 0.3)$$

$$= 150 + 40 - 180 = 10$$

All that remains is to calculate the value of the decision node — by comparing the expected values of the alternatives, and deciding which one is better. The expected value of CD paying 5% is \$50, while the expected value of Risky Investment is \$10. The CD offers the higher expected value and so it is optimal.

When a tree is rolled (or folded) back, the expected value of each node is drawn in a box to the right of the node. At decision nodes, the branches of non-optimal alternatives are identified by two slash marks.



The Decision Tree Tutorial Chapter provides detailed instructions on how to use TreeAge Pro to build and calculate the simple investment decision tree.

2.4 Influence diagrams

TreeAge Pro allows you to create influence diagrams. However, influence diagrams cannot be used for model analysis. Refer to the Influence Diagram Chapter for help.

2.5 Further reading

For additional background on decision analysis in general, here are some suggested references:

- *Making Hard Decisions*, Clemen (1996), Wadsworth.
- *Decision Making and Forecasting*, Marshall and Oliver (1995), McGraw-Hill, Inc.
- *Decision Analysis: Introductory Lectures on Choices under Uncertainty*, Raiffa (1968), Random House.

There are also numerous books and journals dealing with the application of decision analysis in specific fields and industries. Here are just a few selected references:

- *Decision Making in Health and Medicine*, Hunink, and Glasziou (2001), Cambridge University.
- *Decision Analysis for Petroleum Exploration*, 2nd Ed., Newendorp and Schuyler (2000), Planning Press.
- *Coping with Risk in Agriculture*, Hardaker, Huirne, and Anderson (1998), CAB International.

- *Introduction to Decision Analysis: A Practitioner's Guide to Improving Decision Quality*, Skinner (1996), Probabilistic Publishing.

2.6 Example Models

The TreeAge Pro installation includes numerous tutorial example trees, including some examples from the above texts.

3. TreeAge Pro Interface

TreeAge Pro uses a modern, flexible user interface consisting of editors and views.

The interface includes:

- Flexible modeling workspace with views to access different application features.
- Customizable perspectives to show collections of views related to modeling, analysis, debugging or other sets of tasks.
- Tree diagram editor that supports drag and drop functionality for adding/rearranging nodes and multiple editor windows for a single model.
- List filtering to help narrow down lists of variables, tree preferences, etc.
- Auto-completion of reward, probability, variable definition and other expressions.
- Project management tools for organizing models and documents including version control.
- Sharper and more customizable graphical output.
- ... and much more.

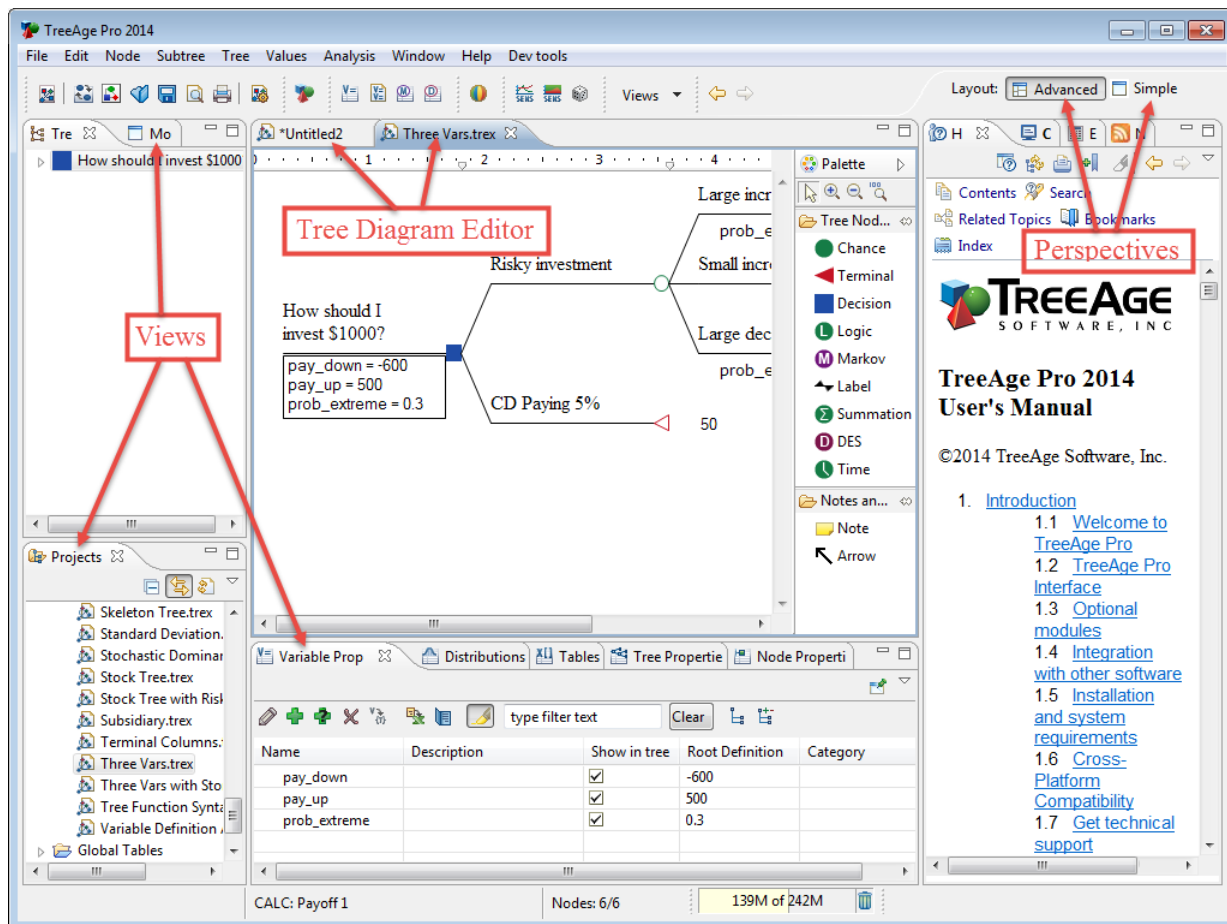
Interface Features

This chapter highlights components of the TreeAge Pro User Interface. Most sections of this chapter can be opened via Online Help within TreeAge Pro.

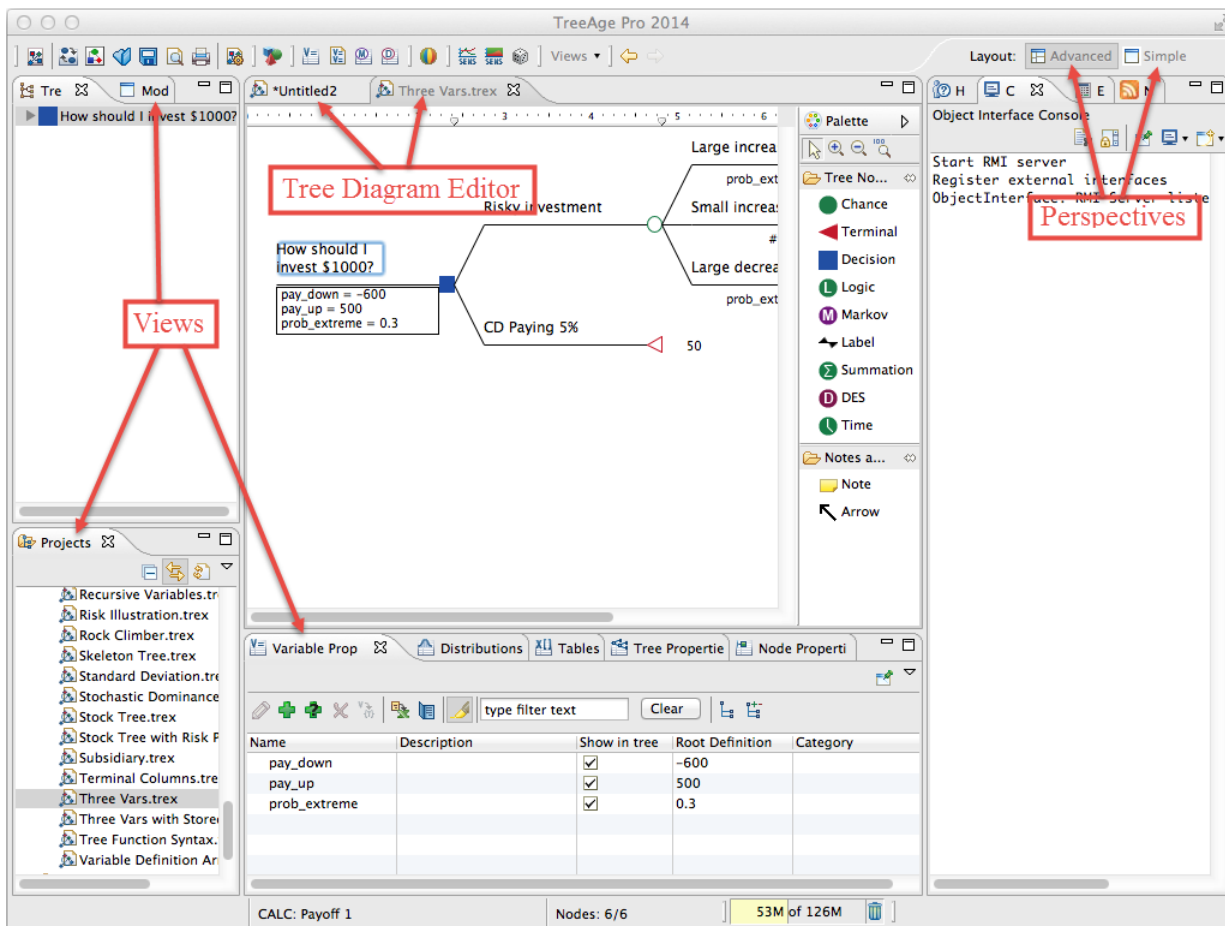
3.1 TreeAge Pro Workbench

The new user interface is called the TreeAge Pro Workbench. The workbench includes a menu and a collection of editors and views that can be resized, collapsed, closed and reopened.

Below are images of the TreeAge Pro workbench with specific user interface elements highlighted.



The TreeAge Pro Workbench - Windows REPLACE ME



The TreeAge Pro Workbench - Mac REPLACE ME

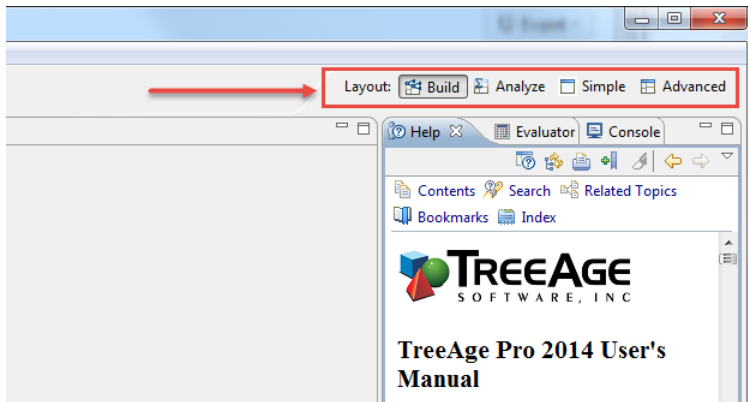
3.2 Perspectives

A perspective (or layout) is a collection of views and editors that makeup the layout of your workbench. Perspectives are linked to the TreeAge Pro workbench and not to a specific document. You can create/use different perspectives associated with different sets of tasks. Such tasks can include modeling, Markov input, running analyses, debugging, etc. Simple, Build and Analysis perspectives come preinstalled with the software.



You may see different perspectives in your workbench based on a workspace created with an older version of TreeAge Pro. To use the new perspectives, you can create a new workspace. Click [here](#) for instructions in the TreeAge Software Knowledge Base.

Perspectives can be selected from the top-right corner of the TreeAge Pro application window. Click on a perspective to change the orientation of editors and views in your Workspace.



Perspectives

As you change the organization of views and editors, the current perspective remembers the layout. The next time you open TreeAge Pro with that perspective, the organization will be the same as when you left it. However, you can return to the original saved version of the perspective by right-clicking on the perspective and selecting Reset from the context menu. You can save the perspective by choosing Window > Save Perspective from the menu. You can then save this new perspective under an existing or a new name. Once the perspective is saved, the Reset option will return the perspective to its new saved organization.

You can customize the active perspective's menus and toolbars via the Customize option from the right-click context menu.

You can close a perspective via the Close option from the right-click context menu. This will remove the visible button for that perspective from the workbench. You can show hidden perspectives by choosing Window > Show Perspective from the menu; then select the perspective you want to show from the Open Perspective dialog.

The Build and Analysis perspectives show different tools and views to assist with the tasks of building and analysing models respectively. The Simple perspective is a more concise perspective. The three perspectives are shown in the figures below with different tools associated with each. Use can roll over each tool icon with a mouse and the tool tips (comments associated with that tool) will describe what each tool does.

Example02-Variables.trex

Standard treatment: Radiation

New treatment: Surgery and radiation

Evaluate new treatment

cFollowupAnnual = 2K
 cRadiation = 30K
 cSurgery = 50K
 effEradicated = 10
 effNotEradicated = 3
 pEradicateRad = 0.6
 pEradicateRadSurg = 0.8

Radiation eradicates tumor
 pEradicateRad
 Radiation fails to eradicate tumor
 #
 Surgery and radiation eradicate tumor
 pEradicateRadSurg
 Surgery and radiation fail to eradicate tumor
 #

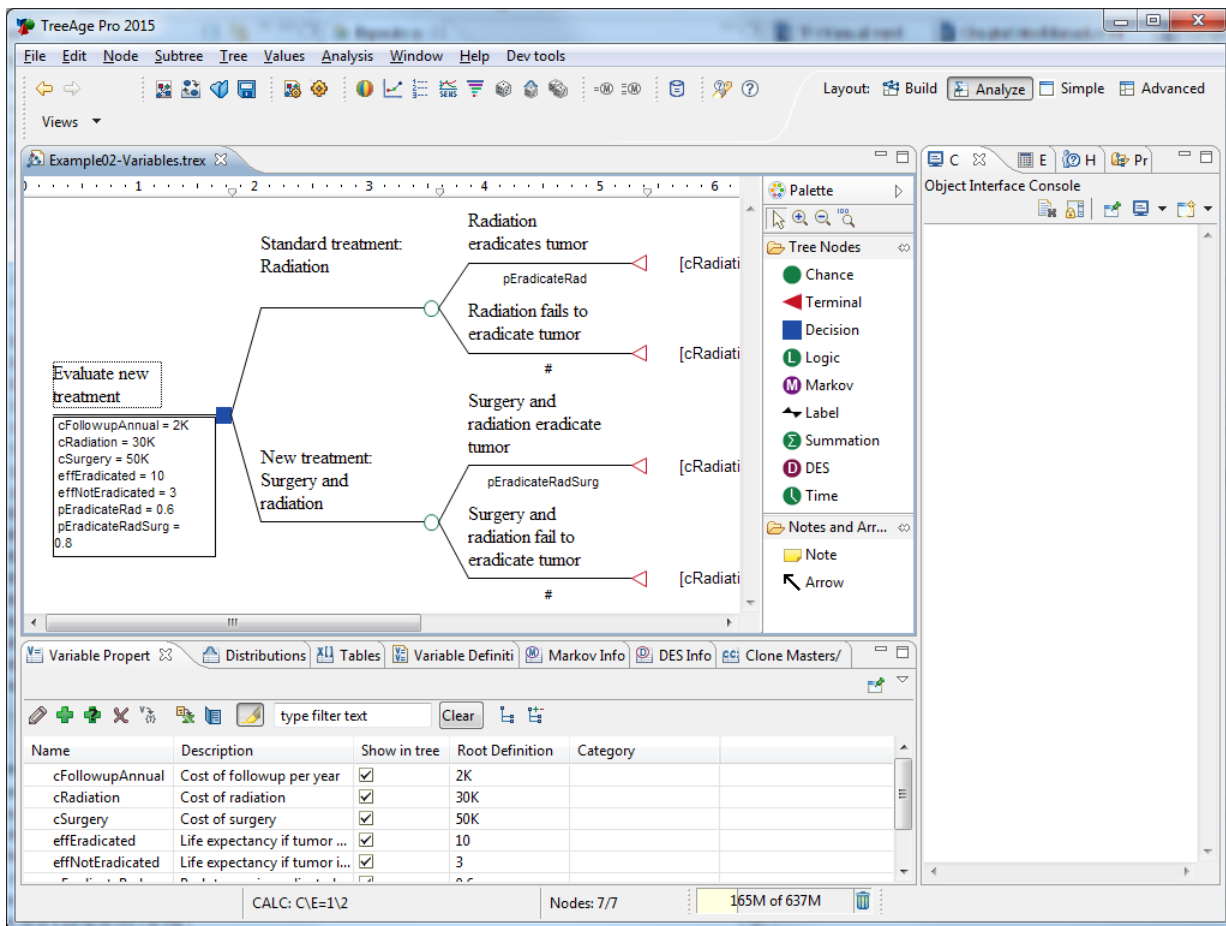
Name	Description	Show in tree	Root Definition	Category
cFollowupAnnual	Cost of followup per year	<input checked="" type="checkbox"/>	2K	
cRadiation	Cost of radiation	<input checked="" type="checkbox"/>	30K	
cSurgery	Cost of surgery	<input checked="" type="checkbox"/>	50K	
effEradicated	Life expectancy if tumor ...	<input checked="" type="checkbox"/>	10	
effNotEradicated	Life expectancy if tumor i...	<input checked="" type="checkbox"/>	3	
pEradicateRad	Prob tumor is eradicated ...	<input checked="" type="checkbox"/>	0.6	

CALC: C/E=1\2
 Nodes: 7/7
 144M of 637M

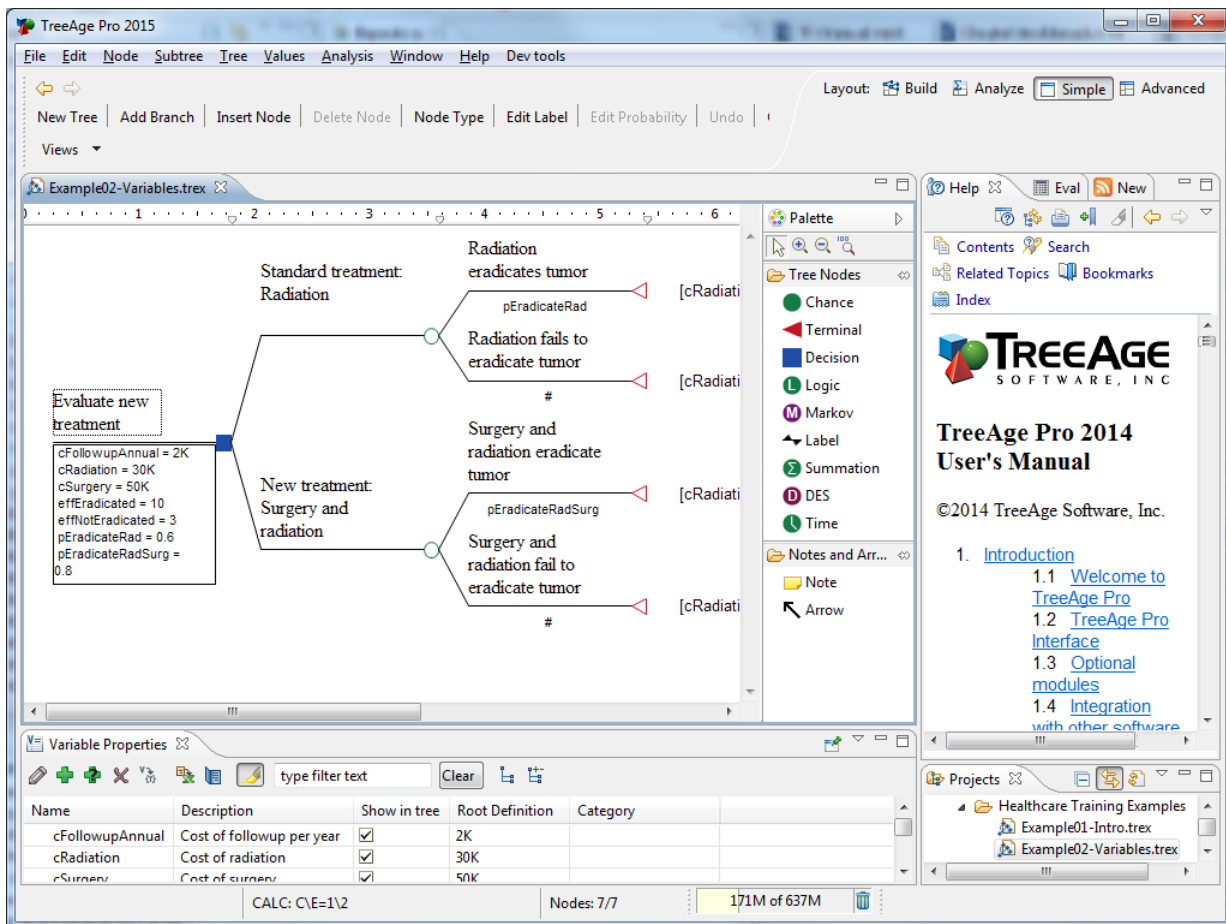
TreeAge
 SOFTWARE, INC.
TreeAge Pro 2014 User's Manual
 ©2014 TreeAge Software, Inc.
 1. [Introduction](#)
 1.1 [Welcome to TreeAge Pro](#)
 1.2 [TreeAge Pro Interface](#)
 1.3 [Optional modules](#)
 1.4 [Integration with other](#)

Project
 Healthcare Training Examples
 Example01-Intro.trex
 Example02-Variables.trex
 Example03-Clones.trex
 Example04-Dominance.trex
 Example05-DrugComparison.trex
 Example06-PSA.trex
 Example07-MarkovSimple.trex

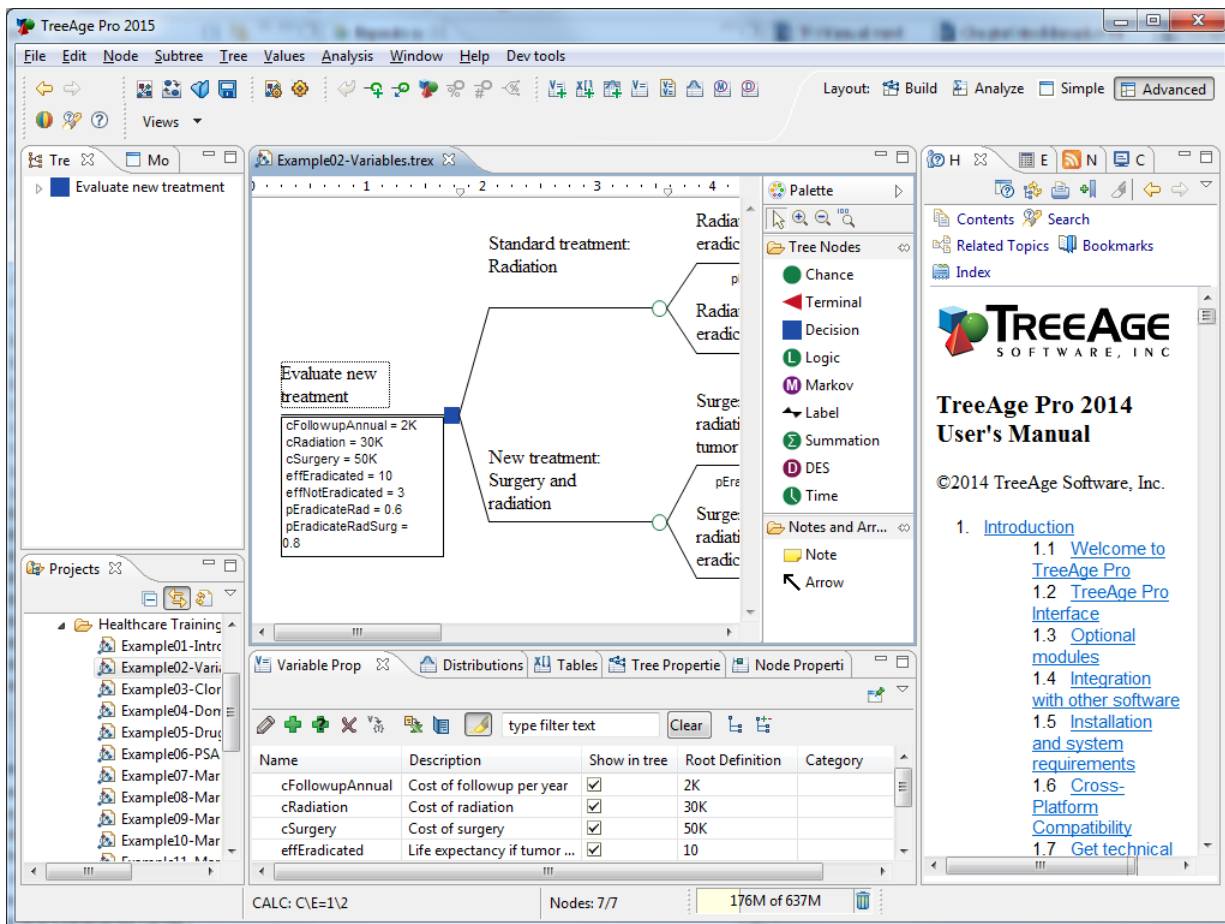
Build perspective



Analyze perspective



Simple perspective

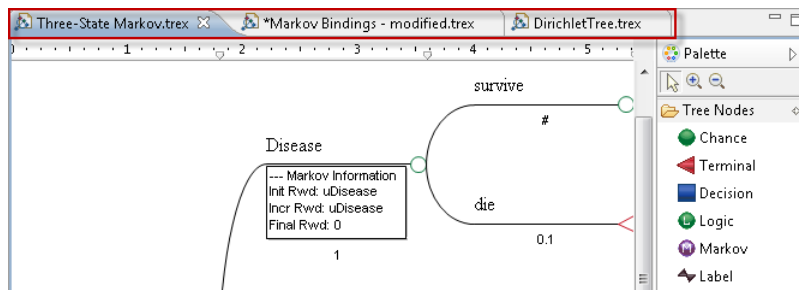


Advanced perspective

3.3 Tree Diagram Editor

The Tree Diagram Editor is the primary modeling window within the TreeAge Pro workbench. You create the model structure within the Tree Diagram Editor by adding, editing, moving and deleting nodes within the model. Changes made within an editor are not saved until you instruct TreeAge Pro to save the model.

You can open multiple Tree Diagram Editors to edit multiple models. You can also open multiple synchronized editors for the same model. When multiple editor windows are open, the inactive windows will appear as tabs at the top of the active window. The active editor window controls the context for all other views in TreeAge Pro.



Tree Diagram Editor Tabs

The Tree Diagram Editor consists of a modeling pane and a palette.

Note that the main application toolbar includes arrows to that allow you to move among the open tabs for models and analysis output.

3.3.1 Modeling Pane

The Modeling Pane is the primary editing window, which contains the visual representation of the model. If the size of the model exceeds the size of the Modeling Pane, scroll bars will appear.



Double click on the tree's editor pane to maximize the Tree Diagram Editor to see more of a large tree.

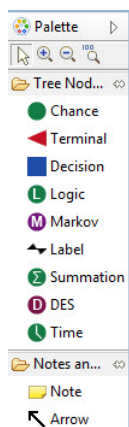
Several actions are performed within the Modeling Pane:

- Click on a node label to enter/edit label text.
- Drag nodes to different places within the model.
- Enter probabilities beneath the branches of chance nodes.
- Select a node to change the perspective of node-specific views (Node Properties, Variable Definitions, etc.)

Modeling Pane edit options

3.3.2 Modeling Palette

The palette contains tools for creating model structure and for viewing the model.



Modeling Palette

Drag nodes of a specific type from the palette into the modeling pane to add a node to the model. The location of the new node is controlled by where it is dropped within the model. The new node line and branch connector will appear as you drag a new node around the existing model.

The zoom in and out buttons allow you to zoom in and out within the modeling pane. First, click on the zoom in or out button in the palette, then click on the model in the modeling pane. The modeling pane will visually recenter and resize based on your selection.

The note and arrow buttons allow you to add those non-structural elements to your model. Click on either element button, then drag the elements onto the model.

3.4 TreeAge Pro Views

TreeAge Pro includes several views to support different features of the software. Almost all TreeAge Pro views are tied to the tree document in the active Tree Diagram Editor window. A subset of the views are further tied to the selected node within the active editor.

The following table lists each TreeAge Pro view, along with a link to the section(s) of the Manual where each view is described in detail.

View/Description	Manual Reference
<i>Model Dashboard:</i> Provides an overview of the important model configuration and model inputs in the model	Model Dashboard and Documentation Chapter
<i>Projects View:</i> Manages sets of files associated with a project, including but not limited to model documents. The Projects View provides a standard tree/explorer interface to help you navigate to files and open files with double-click.	Managing Projects and Documents Chapter
<i>Tree Explorer View:</i> Allows you to navigate to nodes within a tree using a standard tree/explorer navigational interface.	Tools and Functions for Complex Trees Chapter

View/Description	Manual Reference
<i>Model Overview</i> : Presents a miniature view of the active tree in the Tree Diagram Editor. The portion of the model that is currently visible in the Tree Diagram Editor is highlighted.	Tools and Functions for Complex Trees Chapter
<i>Tree Properties View</i> : Edits the properties of a tree. Includes several tabs for different tree elements.	TreeAge Pro 201x Interface Chapter
<i>Variable Properties View</i> : Edits variables in the model.	Working With Variables Chapter
<i>Tracker Properties View</i> : Edits trackers in the model.	Individual-Level Simulation and Markov Models Chapter
<i>Tables View</i> : Edits the properties and data of the model's tables.	Creating and Using Tables Chapter
<i>Distribution Properties View</i> : Edits the model's distributions.	Monte Carlo Simulation, Distributions and Probabilistic Sensitivity Analysis Chapter
<i>Clone Masters/Copies View</i> : Edits the model's clone masters and copies.	Tools and Functions for Complex Trees Chapter
<i>Node Properties View</i> : Edits the properties of the selected node and the elements associated with that node.	TreeAge Pro 201x Interface Chapter
<i>Variable Definitions View</i> : Edits variable definitions of the selected node.	Working With Variables Chapter
<i>Tracker Modifications View</i> : Edits tracker modifications of the selected node.	Individual-Level Simulation and Markov Models Chapter
<i>Markov Info View</i> : Edits Markov info at the selected node.	Markov node: Building and Analyzing Markov Models Chapter Markov state node: Building and Analyzing Markov Models Chapter Markov transition node: Markov Modeling Tools and Techniques Chapter
<i>DES Info View</i> : Edits DES info at the selected node.	DES Models Chapter
<i>State Bindings View</i> : Edits Markov state bindings of the selected node.	Markov Modeling Tools and Techniques Chapter
<i>Evaluator View</i> : Calculates numeric expressions of the selected node.	Working With Variables Chapter
<i>Probability Wheel</i> : Uses the wheel to edit numeric probabilities for branches of the selected node.	Making Changes to Tree Structure Chapter
<i>Console</i> : Views system output.	Calculation Trace Console: Output generated by calculations. Debug output is written to this console.

View/Description	Manual Reference
	Report Console: Output generated by report generation. Object Interface Console: Output generated by the Object Interface.

TreeAge Pro Views

3.4.1 Tree Properties View

The Tree Properties View allows you to edit the properties of a tree. The contents are tied to the model in the active Tree Diagram Editor. Switching editor windows will change the contents of the view.

The following tabs are available within the view to edit particular sets of tree properties:

- General: Edits model document information - creator, description. Also allows you to freeze the model from changes.
- Variables: Edits variable names, properties, default values and sensitivity analysis ranges. The contents of the Variables Tab are the same as those in the Variable Properties View.
- Trackers: Edits tracker names, properties and default values. The contents of the Trackers Tab are the same as those in the Tracker Properties View.
- Tables: Edits table properties and data. The contents of the Tables Tab are the same as those in the Tables View.
- Distributions: Edits distribution properties, type and parameters. The contents of the Distributions Tab are the same as those in the Distributions View.
- Clones: Edits clone master properties. The contents of the Clones Tab are the same as those in the Clone Master/Copies View, which is described in the Tools and Functions for Complex Trees Chapter.

Tree Properties View Tabs

3.4.2 Node Properties View

The Node Properties View allows you to edit the properties of a specific node and the elements associated with that node. The contents are tied to the selected node within the model in the active Tree Diagram Editor. Selecting another node or switching editor windows will change the contents of the view. If no node is selected, the view is disabled.

The following tabs are available within the view to edit particular sets of tree properties:

- General: Edits label, probability (for branch of chance node), and payoffs (for terminal nodes) of the selected node.
- Variables: Edits variable definitions at the selected node. The contents of the Variables Tab are the same as those in the Variable Definitions View.

- Trackers: Edits tracker modifications at the selected node. The contents of the Trackers Tab are the same as those in the Tracker Modifications View.
- Markov: Edits Markov info for the selected node. This tab is only visible for nodes within Markov models. There are three different contexts for the view for Markov nodes, Markov state nodes and Markov transition nodes.
- State Bindings: Edits Markov state bindings at the selected node. The contents of the State Bindings Tab are the same as those in the State Bindings View.

Node Properties View Tabs

Note that variable definitions are associated with a specific node, while variable properties are associated with the entire model.

4. A Decision Tree Tutorial

This chapter is designed to help new users of TreeAge Pro 201x become familiar with the software. It walks you through how to use the software to build decision trees and perform simple calculations.

Suite/Healthcare users: Refer to later chapters for instructions on setting up cost-effectiveness decision trees and on working with Markov nodes and models.

You may want to review the TreeAge Pro Interface Chapter before continuing, in order to familiarize yourself with the many elements of the user interface.

4.1 Constructing a tree

The tutorial in this chapter is based on the investment decision problem described in the Decision Analysis Primer chapter. If necessary, create a new, empty tree now.

4.1.1 Create a new tree

To create a new tree document:

- Select File > New from the menu or click Ctrl-N on the keyboard.

A window will open to prompt you to select the type of template for the new tree document. Select "Blank Tree Diagram" and select "OK".

This will open a new empty tab within the Tree Diagram Editor. The new model will consist of a single Decision Node, which is what we want in this case.



The node furthest to the left within the model is the root node. All models must have a single root node from which all other nodes emanate, either directly or indirectly.

It is best practice to *Enter a label* for the Root node. When you see red text in the Tree Editor, read to text because it is prompting you to enter information.



When you see red text in the Tree Editor, it is prompting you to enter information into your tree. Examples are:

- Enter label: Appears above branches and prompts you to enter a table for the branch.
- Enter prob: Appears below the branches of a chance node and prompts you to enter the probability.
- Enter condition: Appears below the branches of a logic node and prompts you to enter the condition to test at the logic node.
- Enter time: Appears below the branches of a time node branch for Discrete Event Simulation models.

4.1.2 Selecting and deselecting a node

TreeAge Pro shows that a node is selected by filling in the node symbol.

To select a single node using the mouse:

- Click on the node line or the node symbol.

To deselect a single node using the mouse:

- Click in an empty white space anywhere in the Tree Diagram Editor.

4.1.3 Entering node labels

When a single node is selected, a box will appear around the node label text. In the case of a new node, the initial text will be "Enter label".

To enter a node label:

- Click where it says "Enter label", above the node line, and an input text box will appear with a blinking text insertion cursor above the line and to the left of the node.
- Enter a brief phrase in the box to describe the event — in this case, "How should I invest \$1000?".
- Click outside the node to deselect it.

How should I invest
\$1000?



It is generally important to label all nodes in a model to make it clear what each node represents. However, it is possible to put some or all details in note boxes or node/branch comments (which are hidden). Note boxes can either be displayed or hidden, see section 13.3 LINK.



- If you select a single node, you can use the Node Properties View to update the node label.
- When a single text "container" (e.g., a node label or probability, or a note box) is highlighted, the F2 key will activate the text cursor in the selected text part and select existing text. The ENTER key will then deactivate editing, saving changes.
- Note label text will auto wrap as needed. You can use the ruler at the top of the Tree Diagram Editor to resize the length of a branch (and all branches directly above and below it), in order to avoid text wrapping.

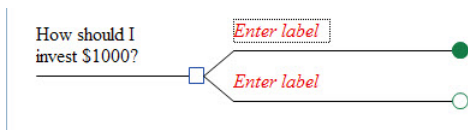
Text Tips

4.1.4 Adding branches/nodes

Branches must be added to the decision node to represent the available investment choices. There are two primary ways to accomplish this task.

To add branches:

- Drag a chance node from the Tree Diagram Editor Palette to the existing decision node. As you drag the new node onto the existing tree, it will appear red in spots where the node can be placed. Release the mouse when the new node is visible as a branch of the decision node.
- Repeat this for a second chance node (also a branch of the decision node).
- ... OR ...
- Right-click on the decision node and select Add Branch from the context menu. Two chance nodes will be added as branches.



Don't be confused by the terms branch and node. They both refer to the same element of a tree model. The term branch is normally used to describe the specific nodes that emanate directly from a parent node (the closest node to the left).

4.1.5 Deleting branches/nodes

If you add an extra branch by mistake, there are a couple of ways to fix the tree.

To delete the node that was just added:

- Choose Edit > Undo from the menu (or Ctrl-Z from the keyboard) to undo the last action.
- ... OR ...
- Select the node.

- Choose Edit > Cut from the menu (or Ctrl-X from the keyboard) to cut the node from the model.
- ... OR ...
- Right-click on the node and choose Cut from the context menu.



Note that nodes that are cut from the model can be pasted to another valid location within the model. This is also true for subtrees.

4.1.6 Zoom in or out

You can zoom in or out of a model via the Modeling Palette within the Tree Diagram Editor.

To zoom in or out:

- Click on the zoom in or zoom out icons (magnifying glass) in the palette toolbar.
- Click on the model in the Tree Diagram Editor to recenter the model with the new zoom perspective.

4.1.7 Navigating the tree using the keyboard

In addition to using the mouse to select a particular node, it is also possible to use the keyboard to change the selected node.

Even if no node is currently selected, there is a keyboard shortcut that will select the root node.

To select the root node:

- Hold down the *CONTROL* key and press the *HOME* key.

To move the selection from the current node to an adjacent node, use the arrow keys. Try using the arrow keys to select the top branch.

To move the selection one node to the right:

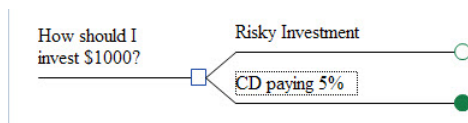
- Press the right arrow key.

4.1.8 Label the strategy nodes

Now, use the actions you have learned to label the strategy nodes.

- Select the top branch of the decision node, and type in its label, "Risky investment".
- Select the bottom branch of the decision node, and name it "CD paying 5%".

The tree should now look like this:





- Rather than entering extensive comments in the node itself, it is often better to use note boxes or node comments (refer to the Annotation Chapter).
- Many tree-building commands are found under the Node menu or by right-clicking on a node.
- If you delete a branch that is a parent (i.e., that has branches), its subtree (i.e., its children as well as their descendants) will not be deleted; instead the children and their descendants will move up one generation. Refer to the Changing the Tree Structure Chapter for more information on inserting and deleting nodes.

Tips

4.2 Entering payoff values

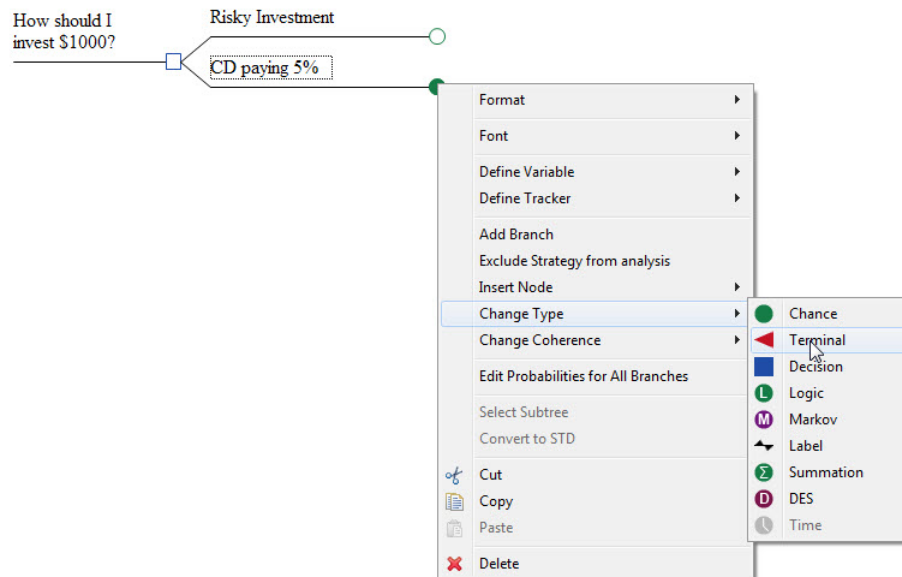
Although the tree structure is not yet complete, it is important to discuss payoffs, which place values on each scenario in the model.

Every path from the root node to an end node within the model represents a scenario. When a scenario is terminated with a terminal node, you must assign value(s) to that scenario. In TreeAge Pro, this is done by entering payoff values.

The *Risky investment* node is not an end point as there are three different possible outcomes associated with that strategy. However, the *CD paying 5%* is the end of a scenario, so it can be terminated.

To change an endpoint to a terminal node:

- Right-click on a node with no branches (in this case the *CD paying 5%* node)
- Select Change Type > Terminal from the context menu (see figure below).
- ... OR ...
- Select a node with no branches.
- Click the "Change node type" icon on the toolbar or click *Control + T* on the keyboard.
- Select terminal from the list of node types.



Change node type via context menu

After changing *CD paying 5%* to a terminal node, TreeAge Pro automatically opens an Enter Payoff dialog for the node.

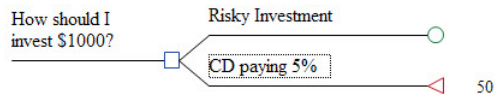
The screenshot shows the 'Edit Payoff - CD paying 5%' dialog box. It has a checkbox 'Use custom payoffs names' which is unchecked. Below it is a table with two columns: 'Payoff Set Label' and 'Value at Node'. The first row is labeled 'Active Payoff' and has the value 'Enter payoff' in the 'Value at Node' column. There is a small icon to the right of the input field. Below the table, it says 'Calculation Method: Simple' and 'Active Payoffs: Active Payoff (1)'. There is an 'Add Payoff Set to Model' button. At the bottom are 'OK' and 'Cancel' buttons.

Edit Payoff Dialog for the selected node

Use the Enter Payoff dialog to assign the numeric payoff for the *CD paying 5%* terminal node. Specifically, set the Active Payoff value to 50 where it says "Enter payoff" and click OK.

Note the "Use custom payoffs names" check box in the Enter Payoff dialogue. If you check this box, you can edit the Payoff Set Label, currently called *Active Payoff* to any name, such as Cost.

The payoff should be displayed in the Tree Diagram Editor to the right of the terminal node, as in the figure below. For more information about setting payoffs see the Payoff Preferences section.



Decision tree with CD paying 5% terminated and payoff set as 50.

To change the payoff values at an existing terminal node:

- Double-click on the payoff values to the right of the terminal node (an elipsis if no payoff values have been entered yet).
- ... OR...
- Right-click on the terminal node and select Edit Payoffs from the context menu.



Further reading:

- The investment decision tree, and most other examples in this manual, requires only Payoff 1. However, each tree can use up to nine different payoffs, or attributes; refer to the Tree Calculation Methods and Preferences Chapter for details.
- See the Healthcare Module documentation, starting with the Building and Analyzing Cost-Effectiveness Models Chapter, for details on setting up cost-effectiveness calculation method trees.
- The Tree Display Preferences and Options Chapter describes how to customize the appearance of payoffs and other visual elements of trees in TreeAge Pro.
- The Introduction to Variables and Sensitivity Analysis Chapter and the Building Formulas Using Variables and Functions Chapter will show you how to use variables and formulas in payoffs.

4.2.1 Complete the structure

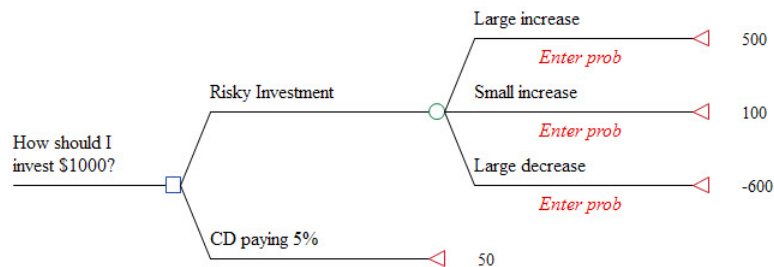
Now that you know how to enter payoffs at terminal nodes, you can finish creating the tree structure. The *Risky investment* strategy is not itself a terminal node, but it needs three branches, each of which is a terminal node.

Add three terminal node branches to *Risky investment* :

- Drag a new terminal node from the Tree Diagram Editor palette into the main editor pane and drop it as a branch of the *Risky investment* node.
- The "edit payoff" dialogue will open automatically. You can enter the payoff value of 500 for Payoff 1. and select OK.Or.....
- If you close the "edit payoff" dialogue before you have completed the details, you can select the prompt in the Decision tree where it says "Enter payoff", and enter the value of 500 and select OK.
- In the tree you will see prompts in red for labels, probabilities and payoffs. Label the branch *Large increase*, by selecting Enter Label above the branch as before.

- This process should be completed twice more, dragging Terminal nodes to the tree, labelling the branches *Small increase* and *Large increase* with Payoff 1 vales of 100 and -600 respectively.

The tree should now look like this:



4.2.2 Saving the tree

Now that all necessary branches have been added, it is a good time to save your work.

Models you create in TreeAge Pro are documents. You save, open, and close trees and other documents in TreeAge Pro the same way you do in other programs — using the File menu commands.

To save a TreeAge Pro document:

- Choose File > Save..., or click the "save" icon (floppy disc) on the tool bar.
- In the Save As dialog, select or create a directory, type "Stock Tree" for the file name, and press enter or click Save.



TreeAge Pro can be set to periodically autosave each document that you open and modify. You can adjust the backup/autosave settings under Window > Application Preferences. In case of a problem, you can recover the latest auto-saved copy. You should still save your work periodically, however. It is also recommended that you save your tree files in a location that is periodically backed up, such as a documents or projects directory on your computer or a network shared drive.

If you open and save files in the Projects View, a local history for the file will be maintained. Right-click on the file in the Projects View to access the local history.

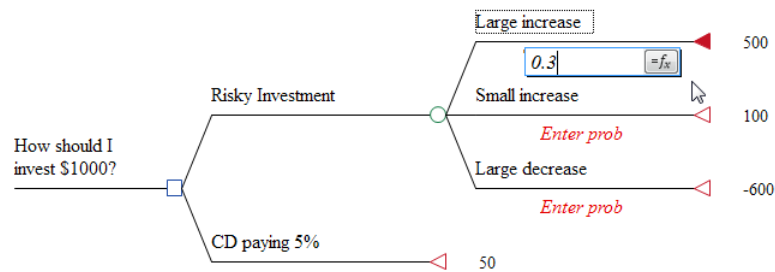
4.3 Entering probabilities

Now, the probabilities must be entered for the three possible outcomes of *Risky investment*. To enter a probability for a chance node's branch, click below the branch where it says "Enter prob". Or, if the branch is already selected, you can edit the probability in the Node Properties View, General Tab.

To enter probabilities:

- Click on "Enter prob" beneath the *Large increase* node. An input field will appear (see picture below).

- Enter 0.3 into the input field and press the enter key.
- Repeat these steps for the *Small increase* probability of 0.4 and the *Large decrease* probability of 0.3.

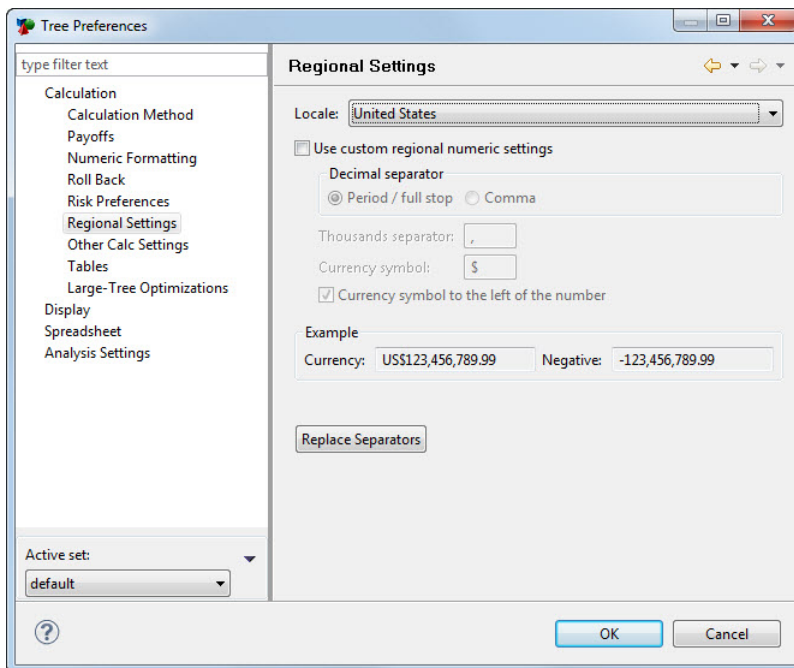


- In the tree diagram, visual cues show where to click to edit probabilities, payoffs, and branch labels.
- Enter label will appear above an unlabeled node to prompt for the label to be entered.
- Enter prob will appear below the branches of a chance node to prompt for the branch probability to be entered.
- Within an input field, function button "=fx" will open an editor where complex expressions using variables and functions can be easily entered.
- Within an input field, the lightbulb indicates that auto-completion is available for this expression. Auto-completion is triggered by pressing CTRL+space on the keyboard.
- The hashmark (#) can be used in place of a probability expression for one branch, in order to have TreeAge Pro automatically calculate the complement during calculations.



Regional (e.g., European) numeric settings:

- If your computer is set up to use commas (",") to represent decimals, rather than periods ("."), you should enter numbers in TreeAge Pro this way. Normally, you will enter numbers in TreeAge models just as you would in a spreadsheet or calculator.
- If your model will be shared by users with different regional numeric settings, use a special tree preference to instruct TreeAge Pro to override a computer's regional settings (i.e., reverse the usage of separators) for that particular tree. This setting is found in the Tree Preferences under Regional Settings (see below).



Tree Preferences, Regional Settings

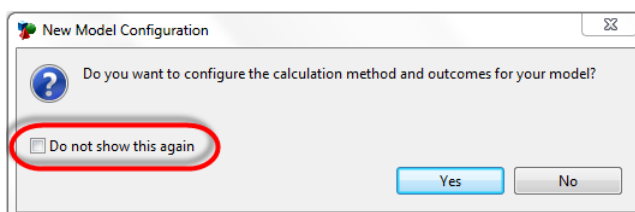
4.4 Configure the model via Tree Preferences

The investment example's tree structure is complete and all values required for calculation have been entered. Before you evaluate the tree, however, a few basic preferences should be specified.

The model preferences can be configured with or without the Configure Model wizard.

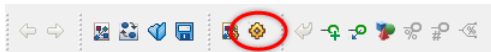
4.4.1 Configuring the model with the Configure Model wizard

When you open a new decision tree, the Configure Model prompt will open and ask you if you want to configure your model. You can tell the software not to show you this prompt again by selecting the check box, as highlighted in the figure below. If you turn this off and want to turn it back on, choose Tree > Auto-Configure New Models from the menu.



Creating a new tree prompts you to use the Configure Model wizard.

If you choose to open the Configure Model wizard at any other time, you can do so via the tool bar, selecting the icon as highlighted in the figure below or by choosing Tree > Configure Model from the menu.

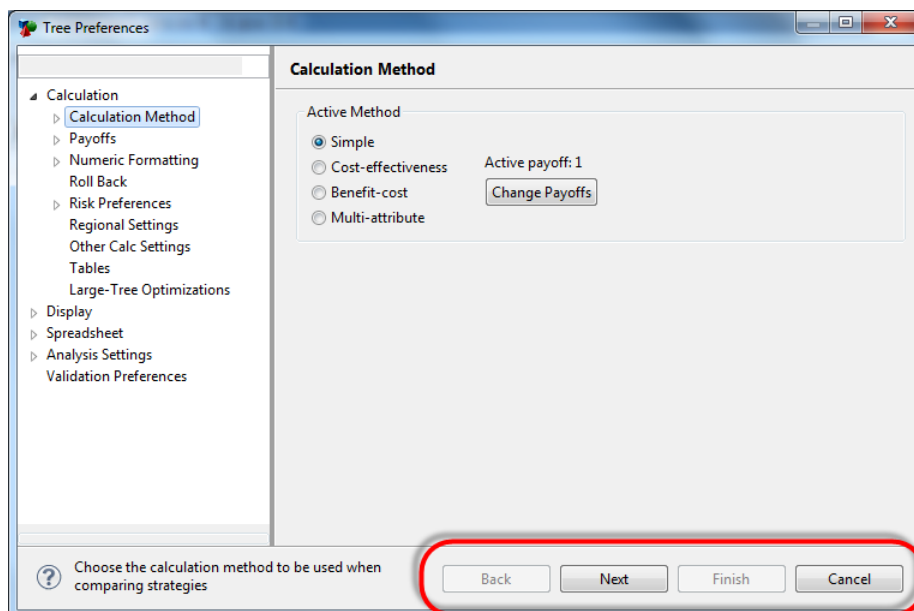


The Configure Model icon, highlighted in the tool bar.

When the Configure Wizard is open, it walks you through the most commonly used the Tree Preferences to allow you to configure the following:

- Calculation method
- Additional options associated with the selected calculation method
- Number of payoffs
- Discounting
- Custom payoff names
- Numeric formatting

The figure below shows the first screen in the Configure Model wizard. You can choose the next, back or finish buttons, as highlighted below, to navigate through the preferences in the model.



The Configure Model wizard walking you through important Tree Preferences.

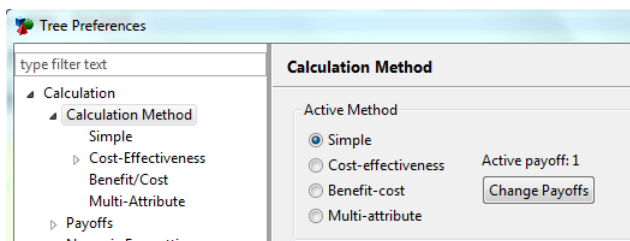
To find out more details about setting preferences, look at the Preferences Chapter.

4.4.2 Configuring the model without the Configure Model wizard

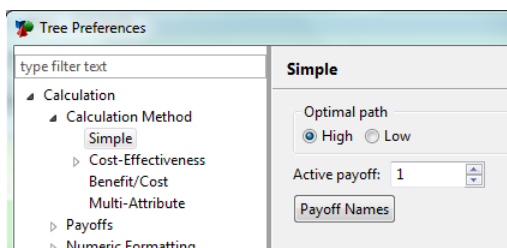
Most importantly, TreeAge Pro needs to know how to select an optimal path at decision nodes. For the investment tree, a strategy that maximizes your income is preferred. You can also specify the appropriate numeric formatting to use when displaying calculated values (number of decimal places, currency symbols, and abbreviations).

To set the calculation method and numeric formatting:

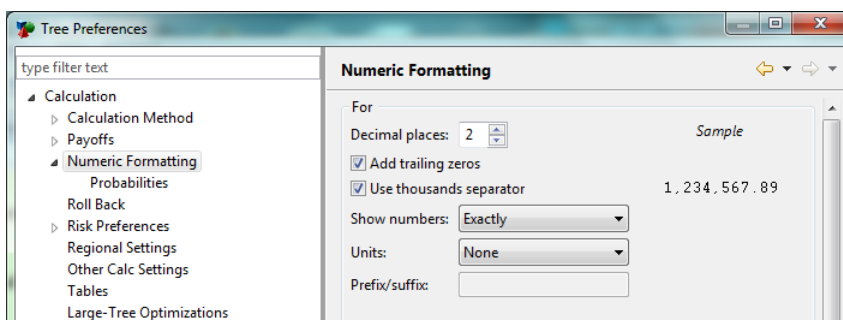
- Choose Tree > Tree Preferences from the menu or press the *F11* key to open the Tree Preferences dialog.
- Select the category Calculation > Calculation Method.
- Select the "Simple" calculation method.
- Select the category Calculation > Calculation Method > Simple.
- Select "High" as the optimal path.
- Leave the active payoff as 1.
- Select the category Calculation > Numeric Formatting.
- Change the Units to "Currency".



Tree Preferences - Calculation > Calculation Method



Tree Preferences - Calculation > Calculation Method > Simple



Tree Preferences - Calculation > Numeric Formatting

Each tree has its own independent set of preferences. Changes to this tree's preferences will not affect other trees.

Press enter or click OK to apply changes, and save the tree again now, by choosing File > Save.

Refer to the Preferences Chapter for more details on numeric formatting and other tree preferences, including information on creating and using preferences sets.

4.5 Calculating the tree

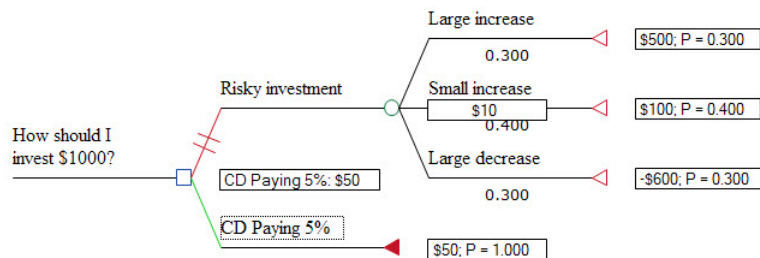
Now the tree should be ready for analysis. Start by rolling back the tree, which performs the expected value calculations described in Chapter 2.

To roll back the tree:

- Select Analysis > Roll Back from the menu.
- ... OR ...
- Click the "beach ball" icon in the toolbar.

The rolled-back tree should look essentially like the picture below.

The optimal strategy, based on the model you have created, will be highlighted by a green branch. Any rejected strategies are highlighted in red with a double "strike through" the arm. If a strategy was excluded from the calculations, this branch would be colored grey.



If TreeAge Pro generates an error message, read it to find out what needs to be fixed. Possible problems include endpoints that are not terminal nodes, and missing probability or payoff values.

If roll back works, but reports look different than those shown below, you may need to check your work, and fix any probability or payoff that was entered incorrectly.

Or, perhaps you have not specified appropriate numeric formatting, as described on the previous page.

Changes cannot be made to structure or values while Roll Back is on (look for a check mark next to the Analysis > Roll Back menu item).

To turn off roll back display:

- Select Analysis > Roll Back from the menu.
- ... OR...
- Click the "beach ball" icon in the toolbar.

4.6 What's next?

This completes the basic decision tree tutorial. You are now ready to use TreeAge Pro to build your own decision trees.

Some questions you may have at this point:

- How can I build a tree to calculate cost-effectiveness? The cost-effectiveness form of multi-attribute calculations is a feature available in the TreeAge Pro Healthcare module. Refer to the Building and Analyzing Cost-Effectiveness Models Chapter.
- How can I include my model in a presentation? Refer to the Printing and Presenting Trees Chapter.
- How can I do sensitivity analysis? Refer to the Introduction to Variables and Sensitivity Analysis Chapter.
- How can I generate a risk profile histogram? Refer to the Analyzing Decision Trees Chapter.
- What if I have very complex cost formulas? Refer to the Building Formulas Using Variables and Functions.
- How can I assign a utility function? Refer to the Utility Functions and Risk Preferences Chapter.

You might take some time now to review the topics in these chapters. As you work with the software and have questions about functionality, bear in mind that the table of contents at the front of the manual and the index at the rear will simplify finding answers.

5. Influence Diagrams

In TreeAge Pro, the primary model type is a decision tree. However, influence diagrams can be helpful when creating or presenting a model. You can create influence diagrams in TreeAge Pro for these purposes. However, all analyses must be performed on decision trees.

This chapter consists of a tutorial describing how to create an influence diagram.



Influence diagrams replace dependency diagrams supported by TreeAge Pro 2012 and earlier.

Influence diagrams provide new functionality as well as all functionality supported by dependency diagrams. Dependency diagrams are obsolete, but they can be imported into the new influence diagram format.

5.1 Influence diagrams

Influence diagrams tend to be simpler on their face than decision trees. While they do not display the level of detail found in a tree (i.e., scenarios, probabilities, and payoffs), influence diagrams portray more clearly the factors to consider in decision making, and how those factors are related. Even in complex problems, where the decision tree is far too large to fit on a single printed page, the corresponding influence diagram is almost always small enough for simple reproduction and efficient communication.

The design of an influence diagram is subject to a number of guidelines. Here are the basic ones:

- *“Nodes” of different shapes represent the factors relevant to the problem.* Each element of the problem — the final objective (e.g., maximizing profit), along with each decision, and random event that can affect the objective — is represented by a single node. A value node (diamond) denotes a measure of the final objective. A decision node (square) is used to indicate a decision. A chance node (circle) is used to represent an event whose value (or outcome) is currently unknown.
- *Related nodes are connected by arcs.* An arc ending in an arrow is drawn between two nodes to indicate that: (a) the first event precedes the second, and/or (b) the first event or action affects (or conditions) the second. An influence arc might indicate that the probabilities for one event depend on the outcome of a prior event or action. An influence arc might also indicate that an action or event makes some contribution to, or deduction from, the final objective (e.g., project cost, or profit).

5.2 Constructing an influence diagram

The tutorial in this chapter explains in detail the software commands needed to build an influence diagram model of the investment decision described in the Decision Analysis Primer Chapter.

To get started, you will need a new influence diagram document.

To create a new influence diagram:

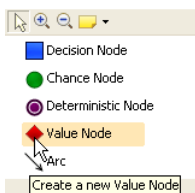
- Choose File > New Influence Diagram from the menu.

5.2.1 Adding and selecting nodes

Unlike a new tree, which starts with a root node, a new influence diagram window is completely blank. The first step in building the diagram is to add the required nodes. Let's start by showing the final objective — profit.

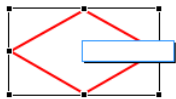
To add a node to an influence diagram:

- Click on the appropriate node type in the Diagram Editor Palette — in this case, the red diamond, for a value node. See below.
- Click and drag the node into a space towards the right side of the influence diagram editor to place it.
- Drag the mouse down and to the right to resize the new node.



Influence Diagram Palette

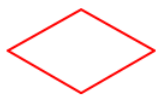
You will see a new, selected value node. TreeAge Pro indicates that the node is selected by showing a rectangular outline with resizing markers on every corner and every edge. The node label will also be selected waiting for you to enter the appropriate text.



New unlabeled node selected

The outline around the node can be used to move or resize the node.

When the node is not selected, the rectangular outline will disappear. Since the node label was not entered, it will appear as an elipsis.



New unlabeled node not selected

To select a node:

- Click inside the node borders.

5.2.2 Entering the node label

You should enter a word or brief phrase in the text box to describe this element of the problem — in this case, the investment objective. You can enter the node label when you first create the node, or you can enter/edit the node label later.

To enter/edit the node label:

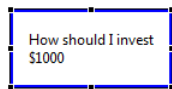
- Select the node.
- Click on the existing node label or the middle of the node if no text has already been entered.
- Type the node label text (in this case *Profit*) in the text area.
- Click outside the node to deselect it.



Profit node with node label entered

Now add a node for the first event that affects return on investment — the decision.

- Create a decision node (blue square).
- For its label, enter *How should I invest \$1000?*



Added decision node



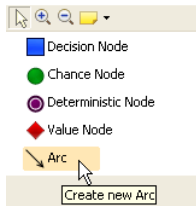
To force a carriage return in a label press *Control + Enter* within the text.

5.2.3 Adding arcs

Earlier in this chapter, arcs were introduced as a means of displaying the relationships between actions, variables, events, and objectives. The direction of influence between the two nodes added so far is from the investment decision to *Profit*, so an arc should be created that points to the value node.

To draw an arc:

- Click on Arc in the Influence Diagram Editor Palette (See figure below).
- Click on the influencing node (*How should I...*) and drag the mouse to the conditioned node (*Profit*); then release the mouse button.



Influence Diagram Palette

A new arc will be created pointing from the *How should I invest \$1000* node to the *Profit* node.



Add new arc

It is frequently useful to add a comment to a label to describe the influence relationship represented by the arc.

To edit an arc label:

- Double-click on the arc.
- Enter text ("Maximize") into the Arc Data dialog Comment field.
- Click OK.

The arc label can also be moved closer/further from the arc.

To move an arc label:

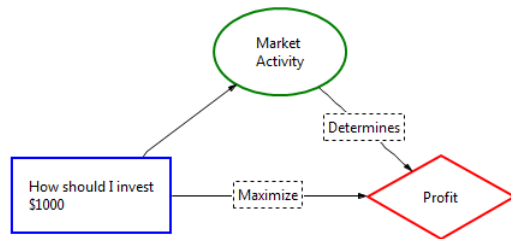
- Click on an arc label and drag it to a new location.

As the arc moves relative to its source/destination nodes, the label will automatically move with it.

Now, add the other required node — a chance node representing the risk inherent in choosing the stock — and its influence arc.

- Create a chance node (green circle), and for the node name enter *Market activity*.
- Create an arc from *Market activity* to *Profit* and enter the label "Determines".
- Create an arc from *How should I invest \$100* to *Market Activity* to indicate the order of actions within the influence diagram.

Here is essentially how your three-node influence diagram should now look:



Influence diagram with third node added

Now that all nodes and arcs have been added, take a moment to save your document. You save, open, and close influence diagrams and other documents in TreeAge Pro the same way you do in other programs — using the File > Save As command. Influence diagrams are saved with the file extension *.indx rather than *.trex.

5.2.4 Outcomes and Alternatives

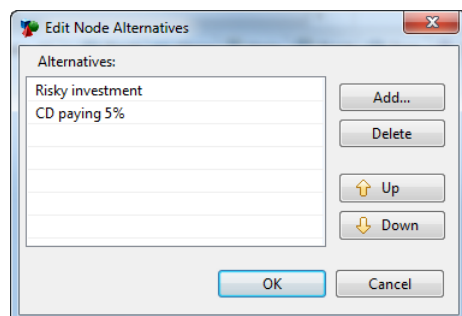
Although the influence diagram may appear to be complete, outcomes and alternatives must be added before it can be converted to a decision tree.

- Alternatives represent the choices available to the decision maker at a decision node.
- Outcomes represent the possible events that can occur from a chance node.

To add alternatives to a decision node:

- Right-click on the decision node.
- Choose Alternatives from the context menu.
- Click Add to create an alternative and enter label text.
- Repeat the last step to add additional alternatives.
- Click OK.

For this influence diagram, the following alternatives should be added.



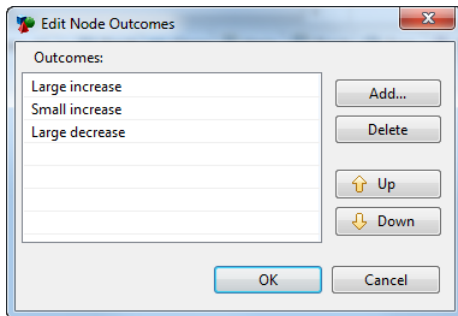
Edit Node Alternatives Dialog

To add outcomes to a chance node:

- Right click on the chance node.
- Choose Outcomes from the context menu.

- Click Add to create an outcome and enter label text.
- Repeat the last step to add additional outcomes.

For this influence diagram, the following outcomes should be added.



Edit Node Outcomes Dialog

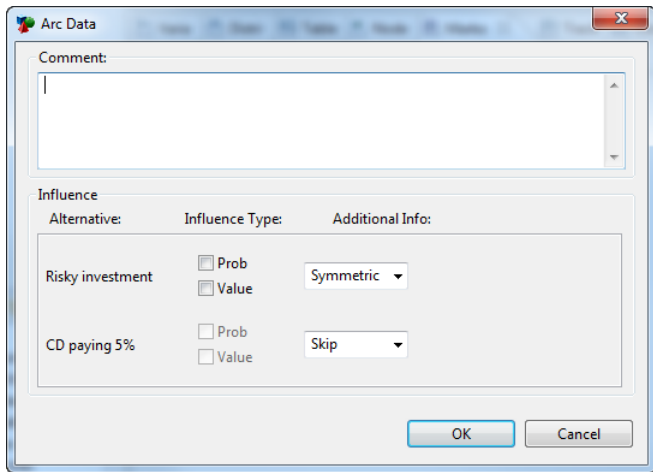
5.2.5 Asymmetry

Asymmetry allows you to eliminate some node combinations from the converted decision tree. For instance, in this chapter's example, market activity affects the *Risky investment* alternative but has no impact on the *CD paying 5%* alternative.

To establish asymmetry in the model:

- Double-click on the arc between *How should I invest \$1000* and *Market Activity*.
- Deselect the Influence Type checkboxes for Prob and Value associated with the *Risky investment* alternative to show that the alternative choice has no influence on the *Market Activity* outcomes.
- Deselect the Influence Type checkboxes for Prob and Value associated with the *CD paying 5%* alternative.
- Choose Skip from the Additional Info column options for *CD paying 5%* alternative to indicate that *Market Activity* has no impact on that alternative.
- Click OK.

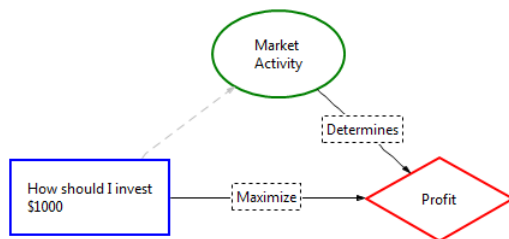
The choices from the steps above are shown below.



Arc Data Dialog

Because there is no impact of the decision alternatives on the chance node outcomes, the arc turns grey and dotted.

The influence diagram is now complete.



Completed Influence Diagram

5.2.6 More editing options

Additional options for editing influence diagrams are described in this section.

To move a node:

- Select a node and drag its outline (not a resize point) to a new location.

When a node is moved, the arcs attached to it will also move.

To cut a node:

- Select a node.
- Choose Edit > Cut from the menu or press *Control + X* on the keyboard. *Arcs attached to the node will also be cut.*

To resize a node:

- Select a node.

- Click and drag one of the resize marks on the rectangular outline around the node.

To select multiple nodes:

- Select one node then hold down the Shift key and select another node.
- ... OR ...
- Click on open space in the diagram and drag to create a rectangle that surrounds all elements you wish to select

To delete an arc:

- Select the arc.
- Press the *Delete* button on the keyboard.

To bend an arc:

- Click on the arc and drag it to another location. Rather than moving the entire arc, it will be split into two line segments, each terminated at the new drag location.
- You can bend this further by dividing up those line segments further using the same technique.

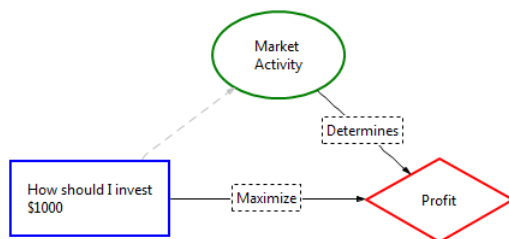
5.3 Converting an influence diagram to a decision tree

Now that the influence diagram is complete, it can be converted to a decision tree.

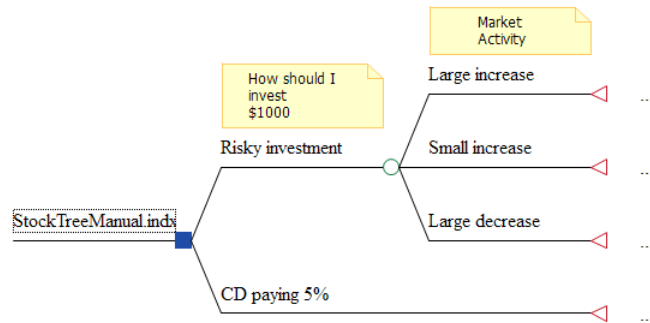
To convert the diagram:

- Open and activate the influence diagram editor window.
- Click the Convert to Tree icon in the toolbar.

A decision tree model is then created in the tree diagram editor with the same name as the influence diagram, but with the extension .trex added to the end.



Influence Diagram

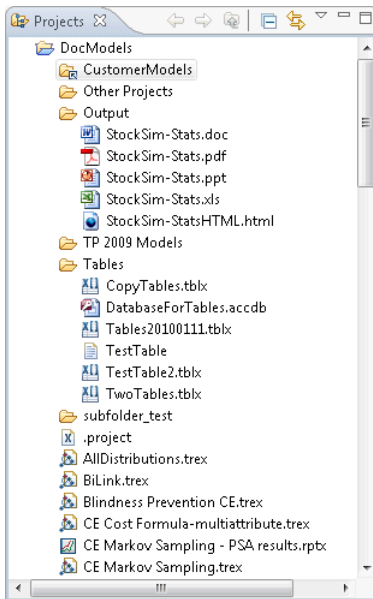


Decision Tree from Conversion

The decision tree structure is created. You would then need to add probabilities and payoffs to the decision tree prior to running analyses.

6. Managing Projects and Documents

TreeAge Pro's Projects View to help organize your model documents.



Projects View

The Projects View provides a hierarchical view of the documents/resources in the TreeAge Pro Workbench. From here, you can open files for editing or select resources for operations such as exporting.

Right-click on any resource in the Project Explorer view to open a pop-up menu that allows you to perform operations such as copying, moving, creating new resources, comparing resources with each other, or performing team operations.



Changes made to your project folders within TreeAge Pro are immediately visible in the Projects View. However, changes to the file system made outside TreeAge Pro are not reflected in the Projects View until you refresh the project via the right-click context menu.

6.1 Create a project

A project is a reference to a folder on the file system on your computer or network where project-related files are stored. At any time, you can create a new project for new or existing models and related documents.

To create a new empty project:

- Right-click in the empty space within the Projects View and select New > Project from the context menu.
- In the New Project Dialog, expand the group General and select Project. Click Next.

- Enter a project name.
- Specify the folder where you want to place the new the project and its files. Uncheck the box "Default location" if necessary.
- You do not need to specify the working sets or referenced projects options.
- Click Finish.

The new project will then appear in the Projects View. Models can then be saved within that project via the menu - File > Save in Project.

To create a new project from an existing folder:

- Right-click in the empty space within the Workspace/Projects View and choose New > Project from the context menu.
- Expand the group General and select Project. Click Next.
- Enter a project name.
- Uncheck the box "Default location".
- Specify the folder where the project files are located.
- You do not need to specify the working sets or referenced projects options.
- Click Finish.

The new project now provides a shortcut directly to the folder (and subfolders) where your project files are located. Existing files in that folder can then be accessed directly from the Projects View.

If a project already exists, perhaps on a network share, you can import it directly into the Projects View.

To import an existing project into your workspace:

- Right-click in the empty space within the Workspace/Projects View and choose Import from the context menu.
- Expand the group General and select Existing Projects into Workspace. Click Next.
- Click on the Browse button next to the Select root directory input.
- Navigate to a folder within which there are one or more projects and click OK.
- The existing projects will be presented in the Projects list. Check the projects you want to select.
- Click the Finish button.

6.2 Tutorial examples project: Example Models

When you start TreeAge Pro for the first time, you will have an Examples project in the Projects View. The Examples project contains the tutorial models that are installed on your computer with TreeAge Pro. Many of these example models are referenced within this manual.

6.2.1 References to Example Models

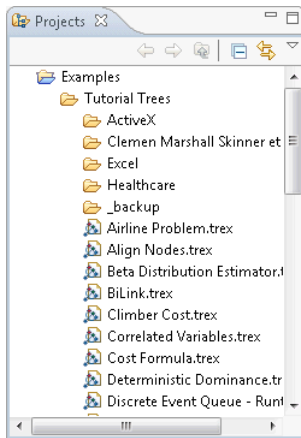
Within this manual, we will reference the Example Models. The following table shows the location of the model we reference them in tutorials. You should refer back to this section often to find the location of the models.

Help file reference to the Example	Where to find the Example in Projects View
Get Started tutorial example	Example Models\ Get Started
Business tutorial example	Example Models \ Business
Health Care tutorial example	Example Models \ Health Care
DES tutorial example	Example Models \ Discrete Event Simulation
Health Care Training tutorial example	Example Models \ Health Care Training Examples
Legal tutorial example	Example Models \ Legal
Oil tutorial example	Example Models \ Oil
Special Features tutorial example	Example Models \ Special Features
Bilinks tutorial example	Example Models \ Special Features \ Bilinks
Clemmens et al tutorial example	Example Models \ Others \ Clemmen Marshall Skinner et al
TreeAge Pro interface tutorial example	Using the TP Object Interface
Other tutorial example	Example Models \ Others

6.2.2 Recreating the Example Models

If, at any time, you delete and modify the Example Models and want to recreate them, you can follow these instructions to do so.

Open the File menu and select Projects > Recreate Tutorial Examples. Follow the prompts, providing a new name for the Example Models. This will create a separate copy of all the example files in addition to those you have modified.



Tutorial Project

6.3 Working within a project

Once you have a project in the Workspace/Projects View, you can immediately open documents within that project via double-click. Double-click will open trees via the Tree Diagram Editor and dependency diagrams within the Diagram Editor. It will also open files that require additional editors like Excel™ and Word™. The right-click menu provides access to file editing options like delete, move, and rename.

In addition, Projects View provides limited version control through the Team, Compare With and Replace With menus. These allow you to compare the current version of the file with prior versions.

6.4 Additional information on Projects View

The Workspace/Project View uses elements from the standard Eclipse Project Explorer View. Click [here](#) for more information on these elements.

6.5 Saving files

When choosing where to save a file, you have two options from the menu:

1. *File > Save in Project*: Save your file within an existing project in the workspace. The file will be saved within the project's root folder, which could be on your computer or on a network.
2. *File > Save As*: Save your file as an independent document on your computer or on a network.

7. Printing and Presenting Trees

This chapter provides basic instructions on how to customize printouts of your TreeAge Pro models, and how to import pictures of models into document and presentation programs like Microsoft® PowerPoint™, Excel™, or Word™.

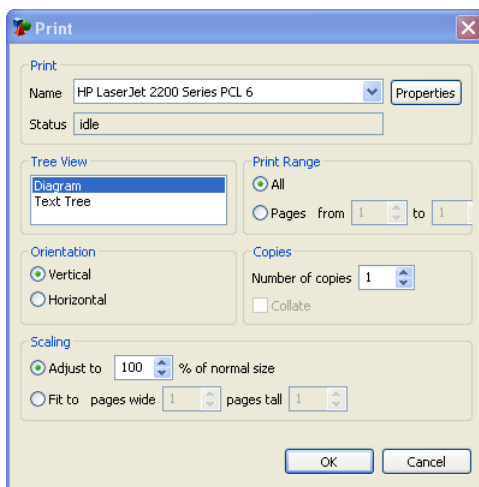
7.1 Printing

Printing documents in TreeAge Pro is similar to printing documents from any other Windows application.

To print a document:

- Select the document you wish to print.
- Choose File > Print from the menu or click on the Print toolbar icon (printer).
- Click OK in the Print Dialog.

The Print Dialog includes some options that may be useful.



Print Dialog

Within the Print Dialog, you can select the Printer from among those available to your computer. The Properties button provides access to the printer's preferences.

The *Tree View* section provides two options.

- *Diagram*: Print the visual representation of the tree.
- *Text Tree*: Print a textual representation of the models nodes based on their position within the tree.

The *Print Range* section allows you to select which pages to print. A small model easily prints on a single page. A large tree may not initially print on a single page, but you may be able to get it to fit by shrinking it, or by changing the page orientation to landscape (horizontal).

The *Orientation* section allows you to select whether to align the paper vertically or horizontally.

The *Copies* section allows you to print multiple copies of the tree.

The *Scaling* section allows you to adjust the size of the tree to fit on more or fewer pages.

You can use Print Preview from the File menu to see how your model will be printed on a page. The Print Preview Dialog has a scaling option to allow you to zoom in/out depending on how the model fits on the page(s).

You can also set page header and footer for the tree via the Tree Preferences.

7.2 Exporting Pictures

You may want to export a picture of a model or part of a model for insertion into a separate document.

To export a picture of a tree:

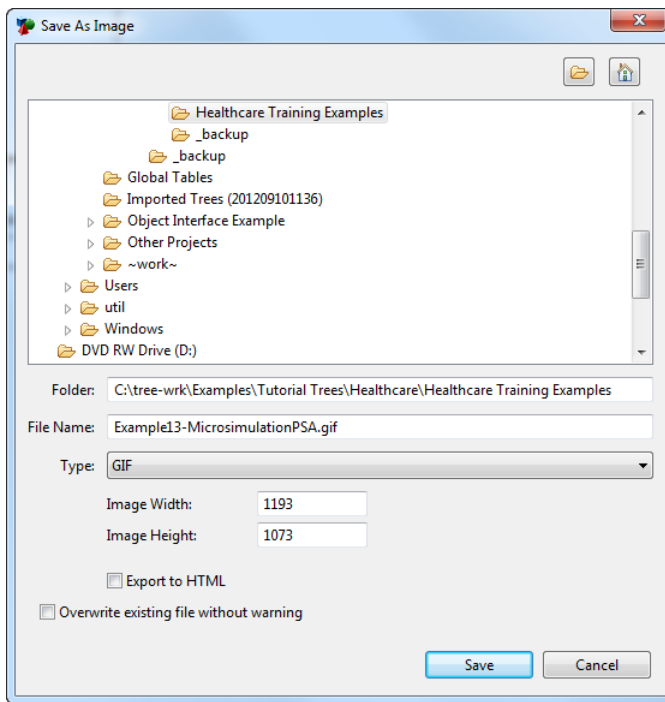
- Select the tree you wish to print.
- From the menu, choose File > Save Image.
- Select the nodes you want to export.

Selected Nodes: Exports the nodes currently selected in the Tree Diagram Editor.

Subtree: Exports the selected node and its entire subtree.

All Nodes: Exports the entire tree.

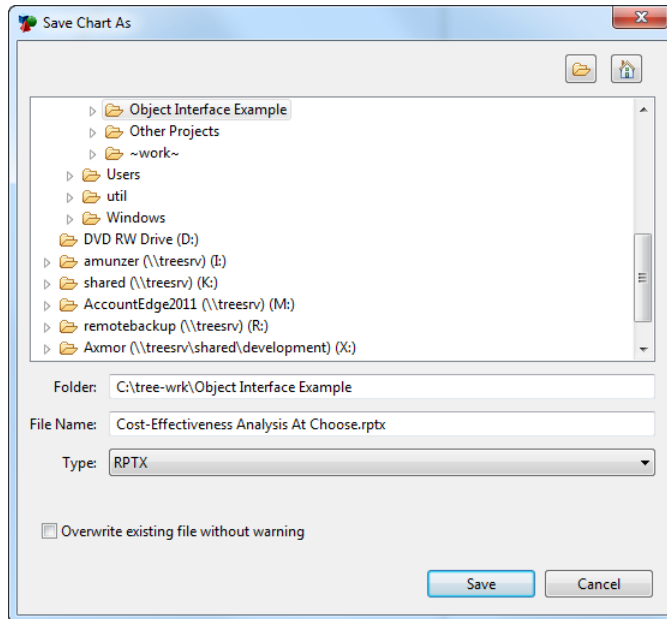
- Select the image format and file location in the Save As Image File dialog box and click OK.



Save As Image File Dialog

To export a graph as an image:

- Create any graph.
- Choose File > Save from the menu.
- Choose Export image file in the Save/Export Chart dialog (see below).
- Enter the filename & location and select the image format in the Save As Image File dialog (see below).
- Click OK.



Save graph dialog



JPEG, JPG and PNG files are good for presentation on the web.

SVG and PDF files are good for sending to publishers or for further editing in graphic design software.

8. Analyzing Decision Trees

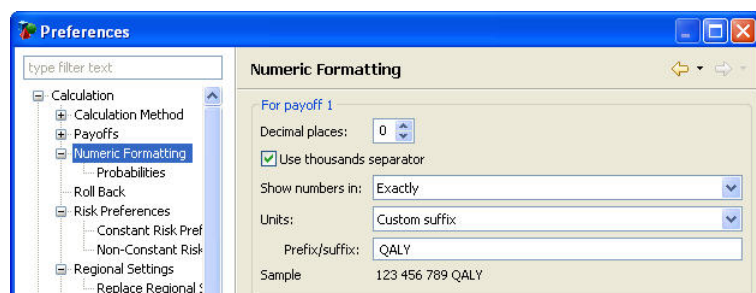
This chapter describes in detail the variety of expected value analyses available in TreeAge Pro.

8.1 Numeric Formatting

TreeAge Pro performs calculations at the highest available precision, but values are displayed using a specified number of decimal places — up to 9 — with the option to use abbreviations and unit symbols. As described at the end of the Decision Tree Tutorial Chapter, each tree and graph has its own set of numeric formatting preferences.

To view/modify a document's numeric formatting:

- Select the model.
- Press the F11 key.
- In the Tree Preferences dialog, select the Numeric Formatting category.



Tree Preferences - Numeric Formatting

Each Numeric Formatting option is described within in the Preferences Chapter.



Notes on entering numbers:

If your computer is set to use commas (",") to represent decimals, rather than periods (".") enter numbers in TreeAge Pro in this fashion, just as you would in a spreadsheet or calculator. Particular trees, however, can be set to override the computer's regional settings and reverse the usage of decimals via the Regional Settings Tree Preferences.

Independent of numeric formatting preferences, payoffs and other values can be entered using K/M/B abbreviations and thousands separators (e.g., typing 2K is equivalent to typing 2,000). Scientific notation can also be used when entering very small or very large numbers (e.g., typing 1e3 is equivalent to typing 1000).

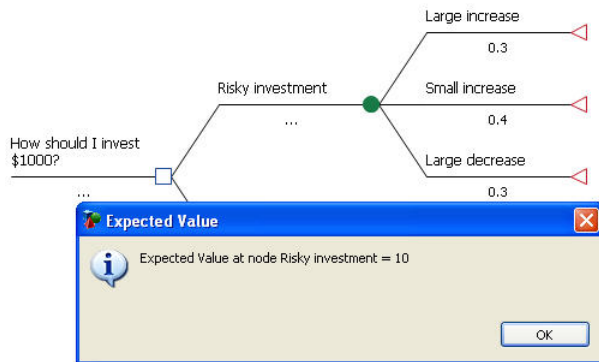
8.2 Expected values

In addition to calculating and displaying expected values for all nodes in a tree (see Roll back in the next section), TreeAge Pro can also report a single expected value for a selected node. This provides a

useful method of verifying the completeness of a single part of an incomplete tree. Roll back calculates and displays expected values for all nodes in a tree (see Roll back, below).

To calculate the expected value of a node:

- Select a node.
- Choose Analysis > Expected Value...
- ... OR ...
- Press CTRL-E on the keyboard.



Expected value at node

The result is displayed using the tree's current numeric formatting preferences; see the previous section on numeric formatting for details.



Using the expected value result:

The Decision Analysis Primer Chapter includes a detailed description of the basic concepts used in calculating expected values in decision trees. Refer to that chapter, or one of the books listed at the end of the chapter for a review of basic concepts.



Calculation details can be reviewed for debugging purposes via Tree Preferences and the Calculation Trace Console. Usage is described in the Markov Microsimulation Chapter, even though the Healthcare Module is not required.

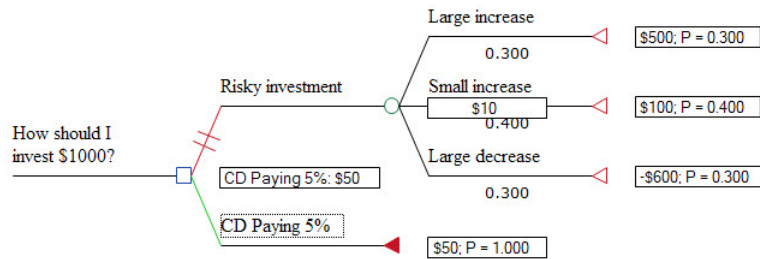
8.3 Roll back

As described in the Decision Analysis Primer Chapter, roll back refers to the calculation of expected values starting at the terminal nodes and continuing back to the root node. In TreeAge Pro, a variety of information is reported in the tree when it is rolled back.

To roll back the tree:

- Select Analysis > Roll Back (or click the beach ball icon in the tool bar).

The rolled back Business tutorial example, Stock Tree, from the aforementioned chapter, is shown below:



Roll back display

Note that expected values are displayed in roll back boxes at every node. The favored strategy, *CD Paying 5%*, is highlighted in green, while the branch connector to the rejected strategy, *Risky investment*, is highlighted in red and displayed with hash marks.

To turn off roll back:

- Select Analysis > Roll Back (again).

8.3.1 Roll back display details

Decision nodes: A box to the right of the node reports the name and expected value of the preferred alternative. TreeAge Pro marks the branches of non-optimal alternatives using hashes, and colors the optimal branch.

Chance nodes: A box to the right of the node reports the expected value. Probabilities are calculated (if necessary) and displayed beneath the node's branches.

Terminal nodes: A box to the right of the node reports the calculated payoff value.

8.3.2 Customizing the roll back display

A picture of the rolled-back tree can be printed or exported to a graphic file; see Printing and Presenting Trees Chapter complete details.

There are many ways to customize the appearance of the rolled-back tree prior to printing or exporting it. In addition to the basic options described on the following page, there are a wide variety of other tree display preferences related to roll back (for example, setting up terminal node columns); see Tree Display Preferences and Options Chapter.

Values displayed in the rolled back are displayed using the tree's numeric formatting settings; see the beginning of this chapter for details on modifying these preferences.

When roll back is turned on, initially all nodes in the optimal path are selected (highlighted). Clicking on the tree will deselect these nodes.

Occasionally, a roll back box will cover the text of a branch description or probability. This can be corrected by moving the box.

To move a roll back box:

- Click and drag the box to a better location.

It is also possible to hide individual roll back boxes.

To hide a selected node's roll back box:

- With the tree rolled back, right-click on the node whose roll back box you want to hide.
- From the pop-up quick menu, choose Hide roll back box.

Notes on roll back boxes:

- Roll back box state will be saved with the tree.
- A hidden roll back box can be redisplayed by right-clicking on the node and choosing Hide roll back box again.
- If you want to reset hidden and moved roll back boxes in a section of the tree, cut the subtree and then paste it back into place.

8.4 Rankings

The Rankings analysis, available when a single decision node is selected, displays a text report listing the alternatives at that node and their expected values, in rank order.

Try the Rankings analysis on the Business tutorial example called Rankings Analysis, which is a slightly more complex version of the Stock Tree from the Decision Tree Tutorial Chapter.

To calculate and rank decision alternatives:

- Select the decision node.
- Choose Analysis > Rankings.

A text report appears which ranks the options, and specifies their expected values. In the case of suboptimal options, it also specifies a marginal (or incremental) value — the amount by which one option is outperformed by the next best option.

The text report dialog includes an Export Report As link to export the displayed text to another program.

Rankings					Actions
rank	name	EV	marginalValue		
1	CD paying 5%	50	0		Export Report As...
2	Blue Chip Stock	32	18		
3	Risky investme...	10	22		

Rankings Analysis Report



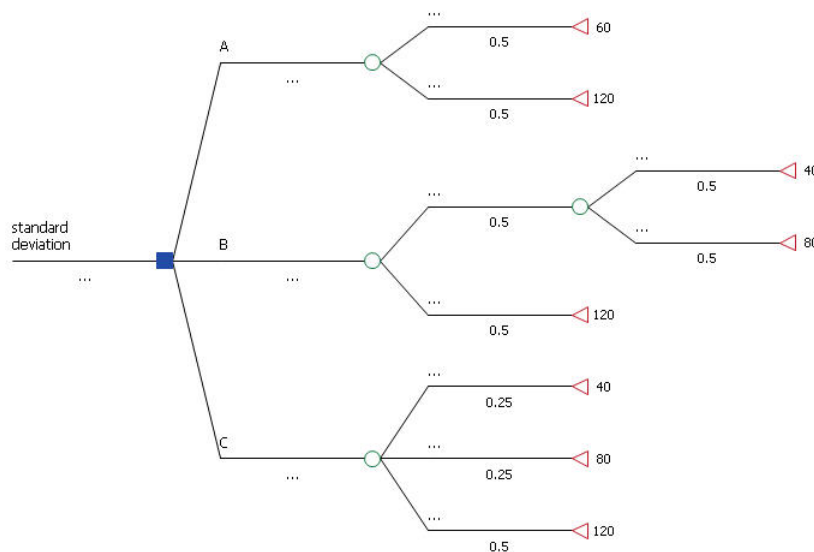
Text Report Tip:

Columns can be resized for easier viewing by clicking and dragging on the dividers between column headings.

8.5 Standard deviation

In addition to comparing strategies based on their expected values, TreeAge Pro also offers several ways to look at an option's risk — the degree of variability in outcomes. The basic statistical measure of risk is standard deviation. In TreeAge Pro, a standard deviation can be calculated for a single strategy (or any other chance node), based on the path probabilities and payoffs of all terminal nodes in its path.

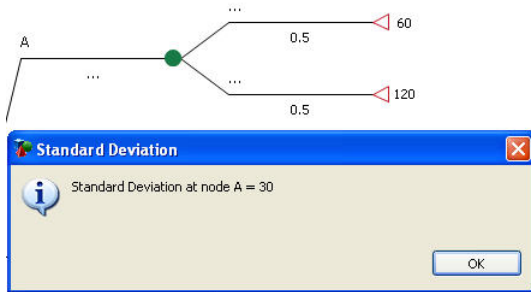
To try the standard deviation calculation, try the Special Features tutorial example, Standard Deviation. In this tree, despite the fact that the three alternatives look different, a rankings or roll back analysis is indifferent between them — all three have the same expected value: 90. In this case, a choice might be based on minimizing risk as measured by standard deviation.



Standard Deviation Model

To calculate a standard deviation:

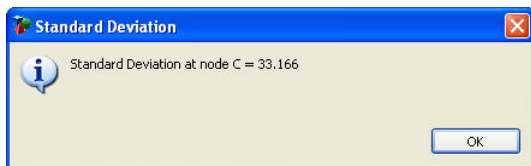
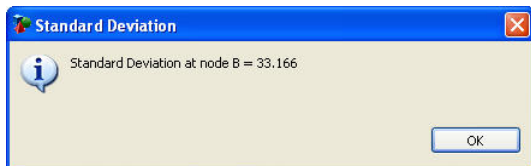
- Select the chance node labeled A.
- Choose Analysis > Standard Deviation.



The calculation used is:

$$SD(\text{Strategy } A) = \sqrt{0.5(60 - 90)^2 + 0.5(120 - 90)^2} = 30$$

where 90 is the strategy A's expected value, or mean. Compare this to the calculated standard deviations for strategies B and C:



If you were to choose a strategy based on minimizing risk, as measured by standard deviation, strategy A would be preferred. Note that B and C are statistically identical, having the same terminal node values and path probabilities, and thus the same standard deviation:

$$SD(\text{Strategy } B/C) = \sqrt{0.25(40 - 90)^2 + 0.25(80 - 90)^2 + 0.5(120 - 90)^2} = 33.166$$

8.6 Discrete simulation/microsimulation

In decision analysis, the most efficient calculation is to use expected values, as described in the Decision Analysis Primer Chapter and illustrated in this chapter. However, it is also possible to evaluate decision trees using simulation, sometimes referred to as discrete simulation or microsimulation.

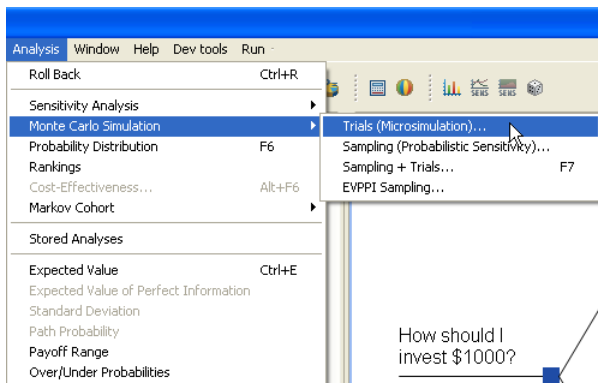
Discrete simulation in decision trees approximates an expected value by “sampling” a representative distribution of paths through the model’s chance events. Discrete simulation of complex models generally utilizes as many “trials” as time allows, in order to improve the EV estimation (ensuring even small probability paths are “sampled” proportionally). If run at a decision node, each trial is repeated for each strategy, to facilitate strategy comparison (e.g., CEA).

- Refer to the chapters on Monte Carlo Simulation and the Individual-Level Simulation for further details on Monte Carlo simulation, including details on running a Monte Carlo probabilistic sensitivity analysis.
- Discrete simulation can also sample values from probability distributions (e.g., in place of a chance node, to represent variability among “individuals”). Refer to the chapter that describes Distributions for the relevant distribution options.
- In survival/Markov models, discrete simulation is particularly useful because it allows modeling any number of continuous and discrete state variables (whereas cohort models work well for more limited numbers of discrete states). The Individual-Level Simulation Chapter provides details on Markov simulation topics.

The current topic provides a very brief example of how to run a discrete (or micro-) simulation in a decision tree, using the simple Stock Tree example from earlier in this chapter. In this model, there are no chance nodes in the “CD” strategy, so there will only be simulation variability in the random walks through the Risky investment strategy.

To perform a simulation (1st-order trials only, no parameter sampling):

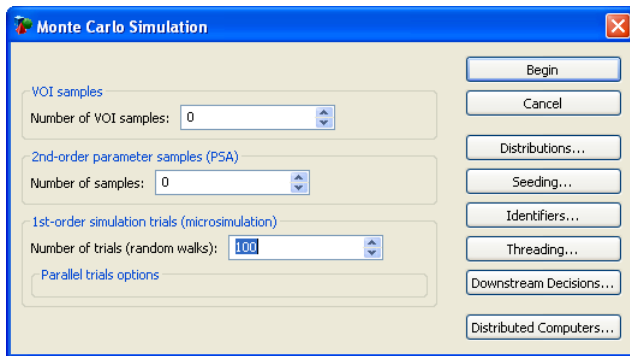
- Select the node for simulation (in the Stock Tree example, the root, decision node).
- Choose Analysis > Monte Carlo Simulation > Trials (Microsimulation)...



- ... OR...
- Click on the dice toolbar button.



- Enter a small number of trials, perhaps 100 and click Begin.



Monte Carlo Simulation Dialog

If the simulation is complex enough to require a significant amount of time to complete, the output window displays the incremental progress of the simulation trials. Once the simulation is done, final statistics and other reporting/graphing options are displayed. Refer to the Individual-Level Simulation Chapter for more details.

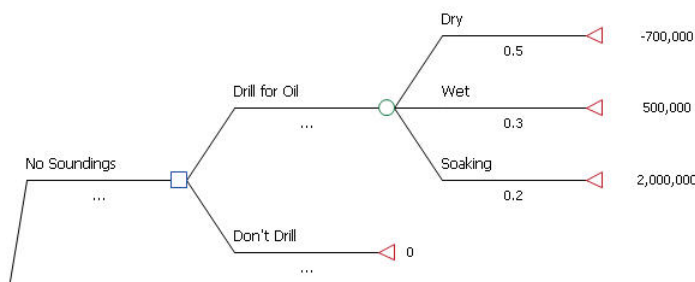
8.7 Probability distributions

The risk associated with alternatives under consideration can be displayed graphically, using a probability distribution histogram, or *risk profile*. A probability distribution graphs the values (i.e., payoffs) and path probabilities of all terminal nodes within a strategy.



This analysis is *deterministic* (no randomness), and does not use the same approach for generating a probability distribution graph as a Monte Carlo simulation (see above). Simulation generates probability distribution graphs based on long runs of *stochastic*, random walks.

Open the Oil tutorial example model, Oil Drilling Problem. This model has some interesting elements, including multiple decisions nodes.



Oil Drilling Problem Model

To create your first probability distributions using TreeAge Pro, start by analyzing a chance node.

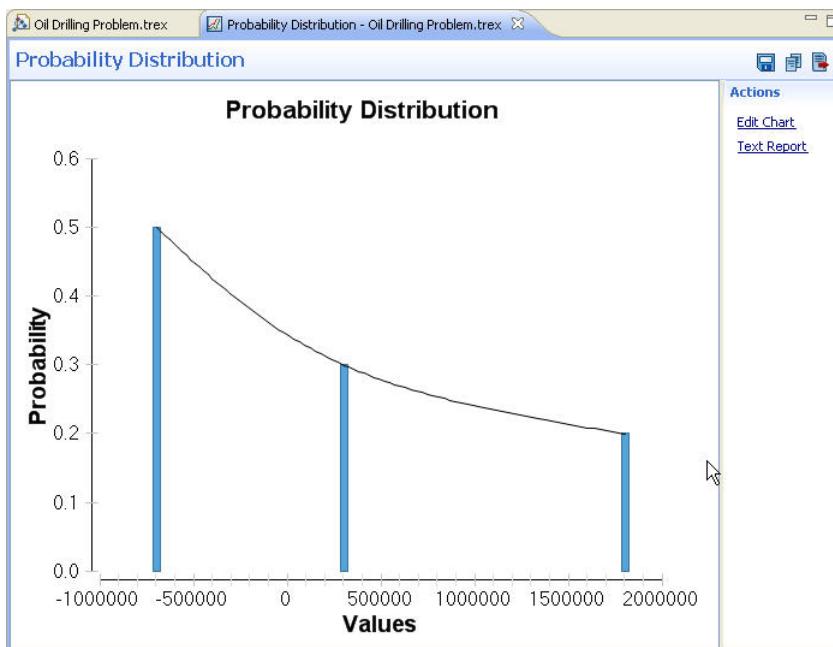
To view a probability distribution histogram:

- Select a chance node – in this case the topmost chance node labeled Drill for Oil, in the No Soundings section of the tree.
- Choose Analysis > Probability Distribution.

TreeAge Pro displays the analysis results in a graph window (see next section). The next section describes some of the options available for graph windows.

8.7.1 Graph window contents

Graphs created in TreeAge Pro are documents that can be printed, saved, or exported to a graphic file. Refer to the Graph Windows Chapter for information on customizing a graph's labels, markers, axes, and other visual elements. Refer to the Excel Output Chapter for information on exporting graphs to Excel.



Probability Distribution Graph

Every graph window includes a set of "Actions" links to the right of the graph. Additional links/controls may be added for special functions, or commonly-used commands. The probability distribution graph includes two links.

- *Edit Chart*: Edit the chart's dimension, scale, format, etc.
- *Text Report*: Show the numerical source data for the graph.

Additional graph options are available via the small toolbar at the top right corner of the graph window. These include functions to save the graph, copy it, or export it. These options are described in the Graph Windows Chapter.

In the example histogram shown in the previous section, TreeAge Pro displays a separate bar for each possible payoff. However, this will not always be the case, as a reasonable number of bars will be used to represent the probability distribution.

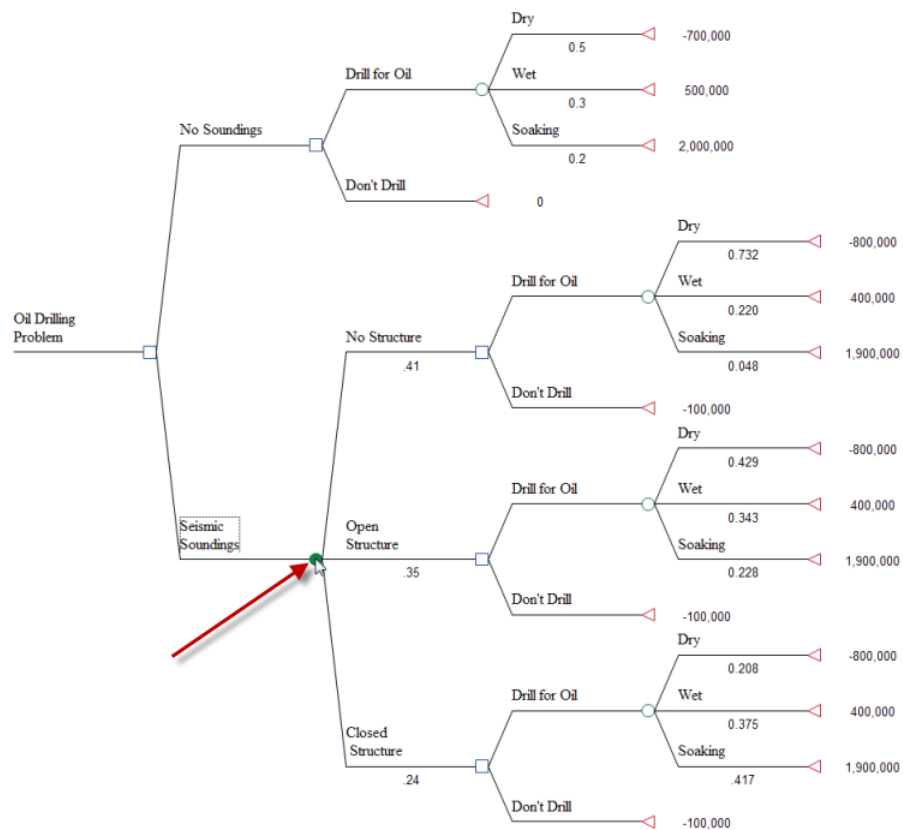
Initially, the value range is divided into a reasonable number of intervals. The height (i.e., probability) of a bar is the sum of the path probabilities of all terminal nodes with payoffs in that interval. The height of all bars sums to 1.0 (100%). The vertical axis will scale to the height of the highest bar. The value/probability associated with each bar can be accessed via the graph's Text Report.

Value	Probability
-700,000	0.5
300,000	0.3
1,800,000	0.2

Probability Distribution Text Report

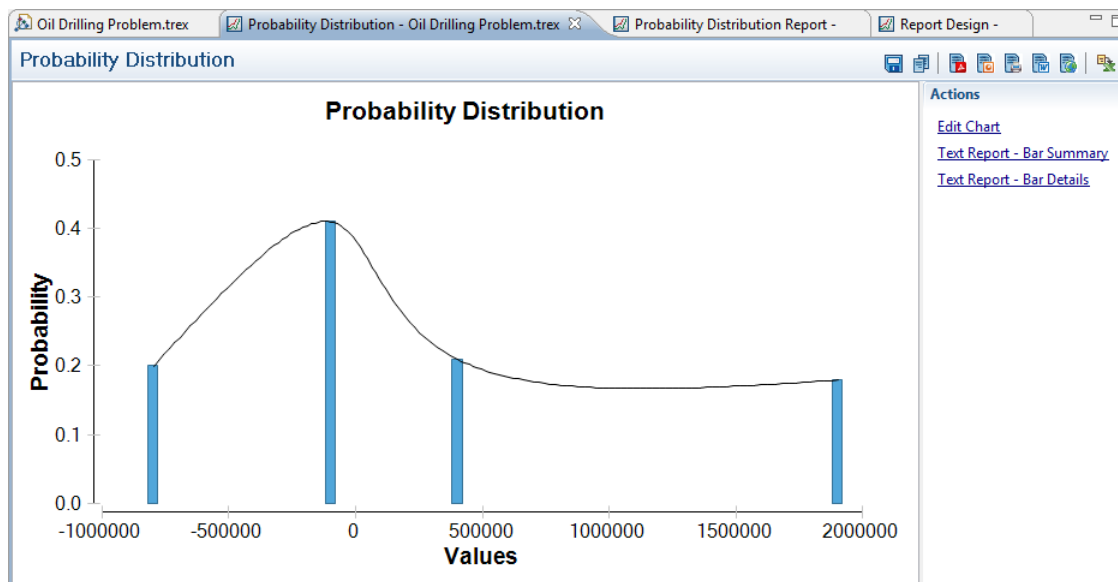
8.7.2 Downstream decision nodes

As noted above, the Oil tutorial example model, Oil Drilling Problem, includes multiple decisions. To see how the probability distribution analysis is affected by downstream decision nodes, analyze the Seismic Soundings chance node. Each of this node's branches is a decision node. See below.



Oil Drilling Problem tree

If there are decision nodes anywhere to the right of the analyzed node, as in this analysis, TreeAge Pro calculates expected values, and then selects an optimal strategy at each downstream, or deferred, decision. The histogram only includes terminal nodes from the optimal path, therefore the path probabilities in the histogram will still sum to 1.0. To see which terminal nodes are in the optimal path, you can roll back the tree.



Probability distribution with downstream decision

The two Text Report links to the right of the graph generate reports showing the underlying data. The Bar Summary report displays the individual bar values and their probabilities. The Bar Details report shows the individual terminal nodes that contribute to each bar. Note that a bar can contain a range of payoff values, even though in this simple example all the payoffs within a bar have the same value.

Probability Distribution

Bar Midpoint (+/-)	Probability
-800,000	0.20007
-100,000	0.41
400,000	0.21005
1,900,000	0.17988

Probability Distribution Text Report - Bar Summary

Bar Midpoint (+/-)	Node	Path Probability	Node EV
-800,000	Dry	0.05	-800,000
-800,000	Dry	0.15	-800,000
-100,000	Don't Drill	0.41	-100,000
400,000	Wet	0.09	400,000
400,000	Wet	0.12	400,000
1,900,000	Soaking	0.1	1,900,000
1,900,000	Soaking	0.08	1,900,000

Probability Distribution Text Report - Bar Details

8.7.3 Cumulative probability distribution

The cumulative probability distribution is not yet supported in Tree Age Pro 201x.

8.7.4 Comparative probability distributions

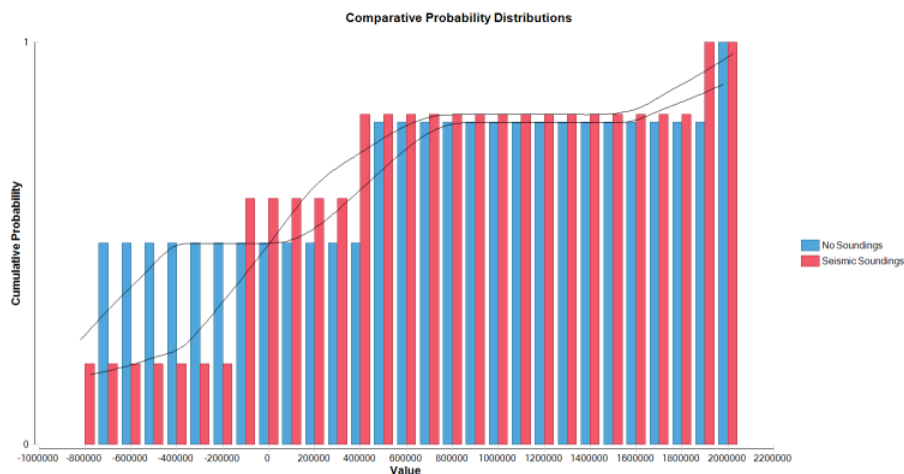
Multiple, cumulative probability distributions can be displayed in a single graph, allowing graphical comparison of options. In a comparative distributions graph, the cumulative distributions are displayed in outline, instead of using filled bars. This format enables graphical comparison of strategies based on their risk profile.

Details about the graphical interpretation of comparative probability distributions are provided later in this chapter, in the section on dominance.

When a single decision node is selected, the Analysis menu displays the Comparative Distributions command. The comparative distributions analysis can be tried at the root, decision node in the Oil Drilling Problem tree.

To generate a comparative probability distribution graph:

- Select a decision node.
- Choose Analysis > Comparative Distributions from the menu.



Comparative Probability Distribution



The figure above was expanded on the screen so that the two strategies' bars did not overlap. Then the screen image was shrunk to fit in this document.

The resulting graph displays the outlines of the cumulative probability distributions for the competing options Seismic Soundings and No Soundings. The outline for a strategy is marked at each corner

(where the cumulative probability “curve” rises) with that strategy’s symbol, as listed in the legend to the right of the graph.

8.7.5 Dominance in probability distributions

Comparative probability distributions can be interpreted graphically, by evaluating conditions of dominance. There are two types of dominance that can be identified relatively easily: deterministic and stochastic (also called absolute and extended dominance). Conditions of dominance can provide more insight into a decision than simple expected value comparison.

Deterministic dominance occurs when one option not only has the best expected value, but its worst possible outcome is better than (or equal to) the best outcome of any other option. It can be identified as follows:

- if optimization requires maximizing value (e.g., profit), the worst “bar” of the dominant option (its left-most vertical line) lies on, or to the right of, the best (right-most) “bar” of the dominated option(s);
- if optimization requires minimizing value (e.g., costs), the worst “bar” of the dominant option (its right-most vertical line) lies on, or to the left of, the best (left-most) “bar” of the dominated option(s).

The kinds of problems in which decision analysis is applied will not often display deterministic dominance, however. Stochastic dominance is more likely. Conditions of stochastic dominance — also called extended or probabilistic dominance — are identified as follows:

- if optimization requires maximizing value, the entire outline describing the stochastically dominant option lies to the right of the dominated option’s outline — the lines can touch for part of the graph, but never cross;
- if optimization requires minimizing value, the outline describing the stochastically dominant option lies on, or to the left of, the dominated option’s outline.

8.8 Expected value of perfect information (EVPI)

Assume that you could buy information that perfectly predicted the outcome of a future uncertainty. What would this information be worth to you?

In a decision tree, the option to acquire perfect information (about a single uncertainty) can be modeled by moving the chance node representing the uncertainty to the left of a decision.

Although the uncertain event still follows the decision in time, the decision maker is assumed to have a perfect predictor of the event outcome before making the decision. Keep in mind that perfect information does not mean that you can control the event’s outcome, only that you can predict the outcome.

Ignoring for the moment the cost of the perfect information, the revised tree cannot have a worse expected value than the original tree, and may have a better expected value. This difference in expected value is referred to as the expected value of perfect information (EVPI).

While predictive information is rarely perfect, the usefulness of EVPI is in calculating a maximum reasonable price for information. If perfect information in a particular situation has a base value of x , one should certainly not pay more than x for imperfect information. To see how imperfect information is dealt with in decision analysis, see the Bayes' Revision Chapter.



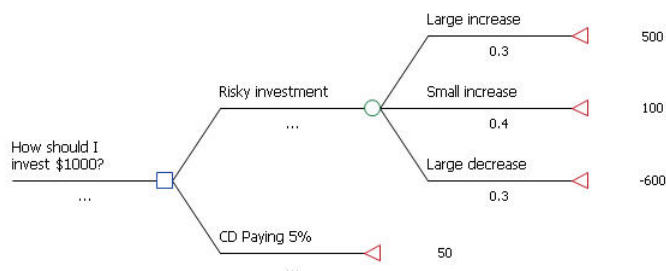
Monte Carlo simulation in TreeAge Pro can be used to calculate expected value of perfect information for any number of predictable or resolvable uncertainties; refer to the Monte Carlo Simulation Chapter.

8.8.1 How EVPI is calculated

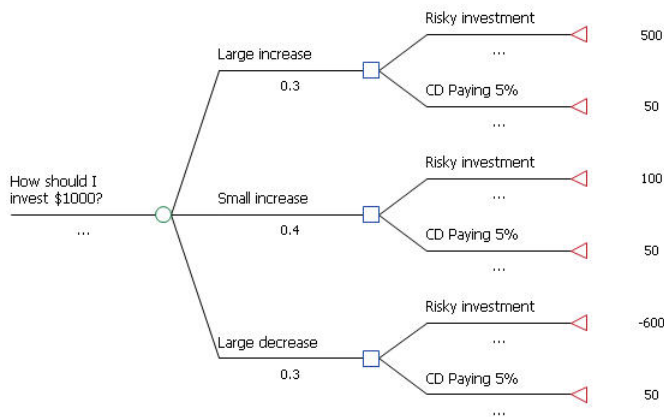
Before trying TreeAge Pro's shortcut for calculating EVPI in a decision tree, it is instructive to work through the extra steps required to calculate EVPI manually.

- Open the Business tutorial example model, Stock Tree.
- Also open the Business tutorial example model, Perfect Information. This version of the investment problem shows the time reversal of the Market uncertainty and the decision.

The two trees are shown below.



Stock Tree



Perfect Information Tree

To calculate the EVPI manually:

- Roll back Stock Tree. The tree's expected value is \$50 (equal to CD paying 5%'s value).
- Roll back the Perfect Information tree. The root node's expected value is \$205.
- To calculate EVPI, take the difference between the expected value of the Stock Tree and that of the Perfect Information tree. The difference is \$155. (If you were minimizing costs rather than maximizing profit, you would subtract the perfect information value from the regular expected value.)

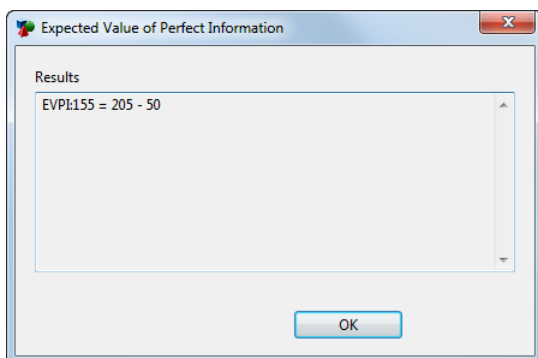
In the investment model, the expected value of having perfect information about the market activity is \$155. This is the most you should be willing to pay to obtain this information, and it affords some basis for appraising to you the value of a less than perfect predictor of market activity.

Now, try TreeAge Pro's shortcut for calculating EVPI in a tree. It requires only the original Stock Tree.

To calculate EVPI automatically:

- Open the Stock Tree.
- Select the Risky investment chance node and choose Analysis > Expected Value of Perfect Info.

A dialog reports the value of \$155.



EVPI dialog

It is also possible to calculate EVPI automatically in a decision tree when the same event appears in more than one strategy, as in example shown below. The assumption in the model is that each of the two stock investments under consideration is followed by the same uncertainty — whether the market will be up or down at the end of the year.

In order to calculate EVPI in this model, both market uncertainty chance nodes must be selected.

To calculate EVPI for the same chance node in multiple paths:

- Before choosing Analysis > Expected Value of Information, *select all nodes in the tree which represent the same event.*

If multiple chance nodes are analyzed, they:

- must be descendants of the same decision node;
- must be “siblings”; and
- must have identical branches using identical probabilities. It does not matter if there are differences in the subtrees further to the right.

If there is more than one decision prior to the selected chance event, TreeAge Pro will prompt you to identify the decision for which EVPI should be calculated.

8.8.2 Avoiding EVPI errors

It is important to understand that it is possible to force invalid EVPI calculations. For example,

- Open the Oil Drilling Problem tree again.
- Select the Drill for Oil node in the No Soundings subtree and choose Analysis > Expected Value of Perfect Info.

In the resulting dialog boxes, you are presented with the option of having the analysis performed at the root, soundings decision node or at the drilling decision.

Performing the calculation at the No Soundings node is similar to the analysis undertaken above in connection with the EVPI tree. It certainly makes sense to calculate the value of knowing the state of oil reserves before deciding whether or not to drill.

However, what if you perform the EVPI calculation at the root decision node? The value reported is \$437,632, significantly higher than at the No Soundings node. Is this a meaningful EVPI calculation?

The structure of the tree already includes the option of securing imperfect information in the form of a seismic test — this is the initial decision. Performing EVPI by placing the chance node representing the uncertain amount of oil to the left of this decision is meaningless. Having already received perfect information, the decision as to whether to obtain additional imperfect information regarding the same subject should have no value or relevance. Refer to the Bayes' Revision Chapter for more information on imperfect information.

8.9 Other Analyses

The last section of this chapter covers three simple analysis tools.

This chapter does not cover all analyses. Refer to later sections of the user's manual for information on additional analysis options:

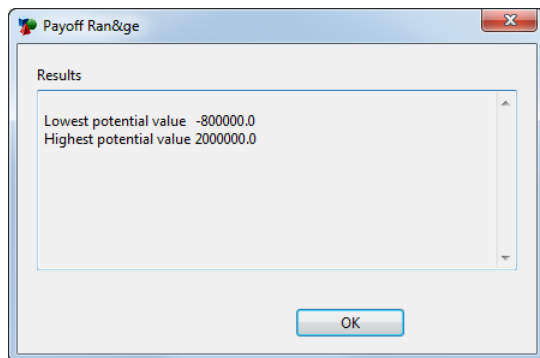
- *Sensitivity analysis* using variables is covered in the Variables and Sensitivity Analysis and More Sensitivity Analysis Tools Chapters. This includes 1-, 2-, and 3-way sensitivity analysis, as well as tornado diagrams.
- *Monte Carlo probabilistic sensitivity analysis* using inputted parameter distributions is covered in Monte Carlo Simulation and Distributions Chapters.
- *Markov cohort analysis* and *microsimulation* are covered in the Building and Analyzing Markov Models and Individual-Level Simulation and Markov Models Chapters.
- *Cost-effectiveness analysis* is covered in the Cost-Effectiveness and Cost-Effectiveness Simulation Chapters.

8.9.1 Range of possible payoffs

This analysis will tell you the highest and lowest payoffs which may occur from the selected node in your tree.

To view the minimum and maximum payoffs in a subtree:

- Select a node, and choose Analysis > Payoff Range.



Payoff Range from root node of Oil Drilling Problem model

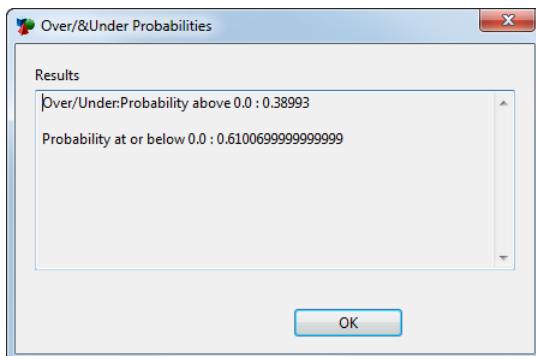
Payoffs from all terminal nodes are included in this analysis, not just those in the optimal path.

8.9.2 Over/under probabilities

The Over/Under analysis calculates the probability of achieving an outcome with a payoff over a target value, and the complementary probability of an outcome under the target.

To calculate the over/under probabilities:

- Select a node and choose Analysis > Over/Under....
- Enter a target value. Indicate whether payoffs equal to the target value should be included in the “under” range. Press enter or click OK.



Over/Under from root node of Oil Drilling Problem model

Over/Under analysis only includes outcomes reached assuming the decision maker follows the optimal path at decisions.

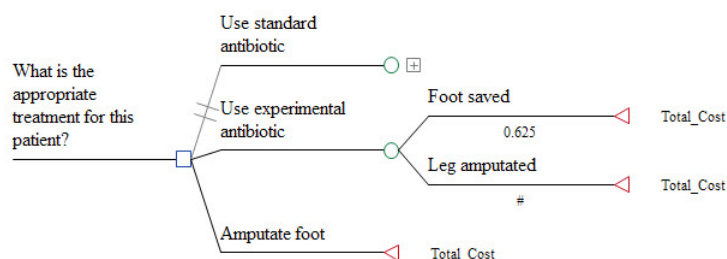
8.10 Excluding Strategies from Analyses

TreeAge Pro allows you to exclude specific strategies from all analyses. When excluded, a strategy will not be presented in the analysis results. The Get Started tutorial example model, Climber Cost, is used to illustrate this technique.

To exclude a strategy:

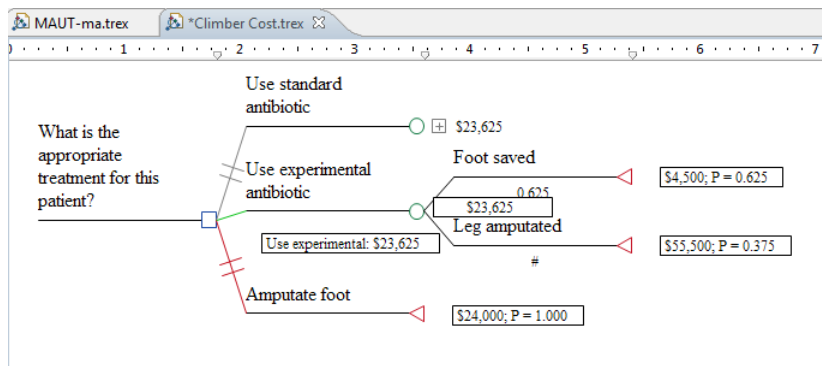
- Right-click on the strategy node you wish to exclude.
- Choose Exclude Strategy from Analysis from the context menu.

The strategy is then marked as excluded by placing hash marks on its branch connector, coloring the branch connector grey and by collapsing the subtree.



Climber Cost model with top strategy excluded, coloring the connector grey.

See how roll back and rankings exclude the strategy.



Roll back with top strategy excluded and the preferred strategy highlighted in green.

Rankings			
Rank	Strategy	Value	Incremental Value
1	Use experimental antibiotic	23625	0
2	Amputate foot	24000	375

Rankings with strategy excluded

Other analyses such as Monte Carlo simulation would also exclude the strategy.

To reverse the strategy exclusion:

- Right-click on the strategy node you wish to exclude.
- Choose Include Strategy in analysis from the context menu.

9. Graph Windows

This chapter includes details on customizing, printing and exporting TreeAge Pro graphs.

9.1 BIRT Project

Beginning with TreeAge Pro 2011, both charts and text reports are generated and displayed using the Business Intelligence and Reporting Tools (BIRT) Eclipse Project. BIRT technology provides highly customizable reporting capabilities for Eclipse-based applications. There are many more options available for customizing charts than in previous versions of TreeAge, as well as a number of new chart "types".

Most users will probably only need to learn a few of the new options and settings for charts/reports that are commonly utilized. This chapter focuses on selected options/features that may be helpful to you.

If you are interested in learning about BIRT capabilities and features in more depth, refer to the printed and/or on-line user's guides written specifically for BIRT users.

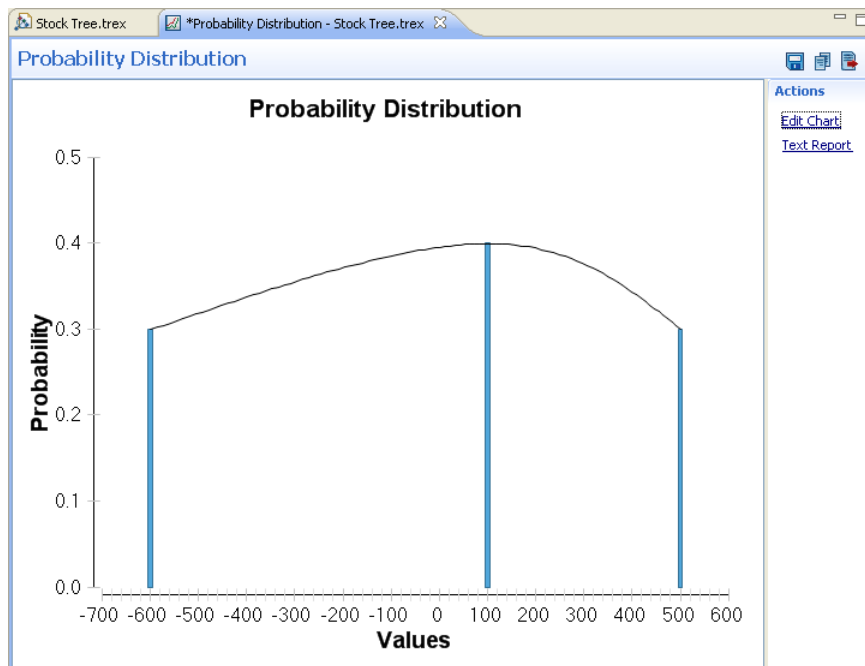
Note that native BIRT functionality has been overridden or disabled by TreeAge Pro in a few cases, in order to support non-standard chart types (e.g., 2-way sensitivity region charts).

9.2 Customizing graphs/charts

Whenever TreeAge Pro displays a graph, you will see two links to the right of the graphical output (specific graph types will provide additional links).

1. *Edit Chart*: Customize the graph's appearance.
2. *Text Report*: Examine the graph's source data.

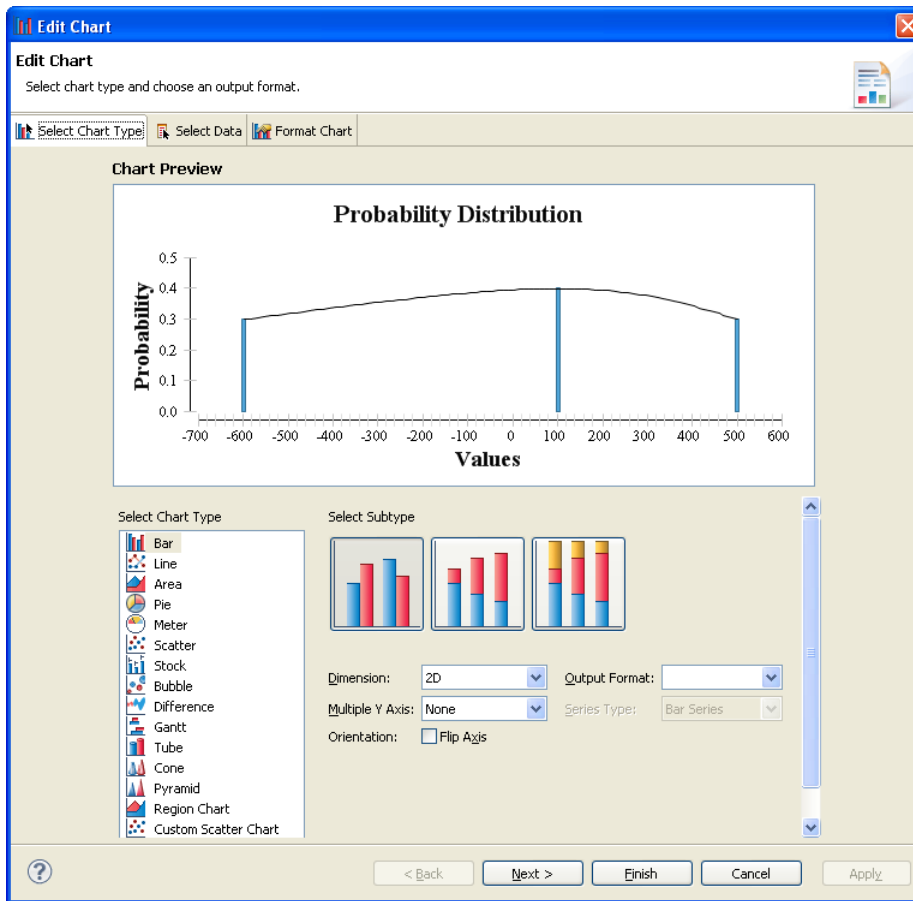
This section focuses on options associated with Edit Chart. The next few subsections focus on general graph customization. We will use the probability distribution graph generated from the *Risky investment* strategy of the Get Started tutorial example tree "Stock Tree" to examine these customizations. However, they are not limited to probability distribution graphs.



Stock tree - probability distribution

9.2.1 Changing the chart type and orientation

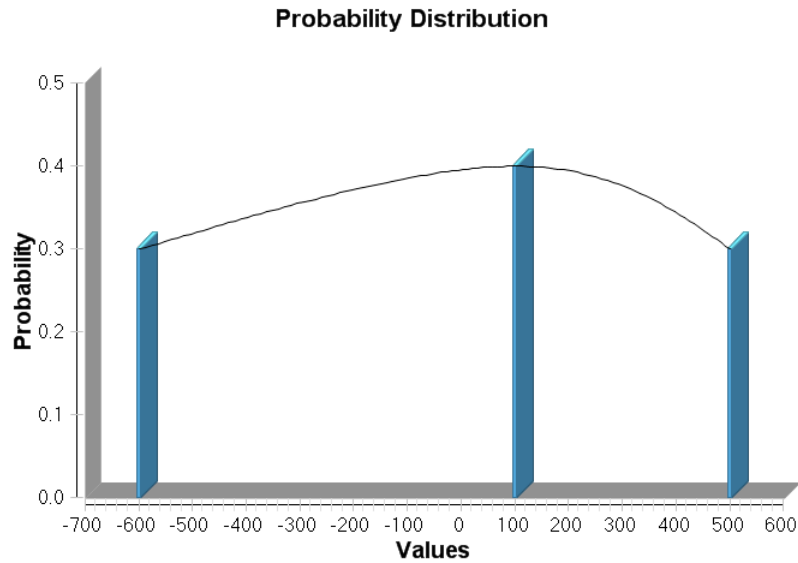
A probability distribution graph is a standard bar graph. If you click the Edit Chart link, an Edit Chart dialog opens providing a large number of options for the graph.



Edit Chart Dialog - Select Chart Type tab

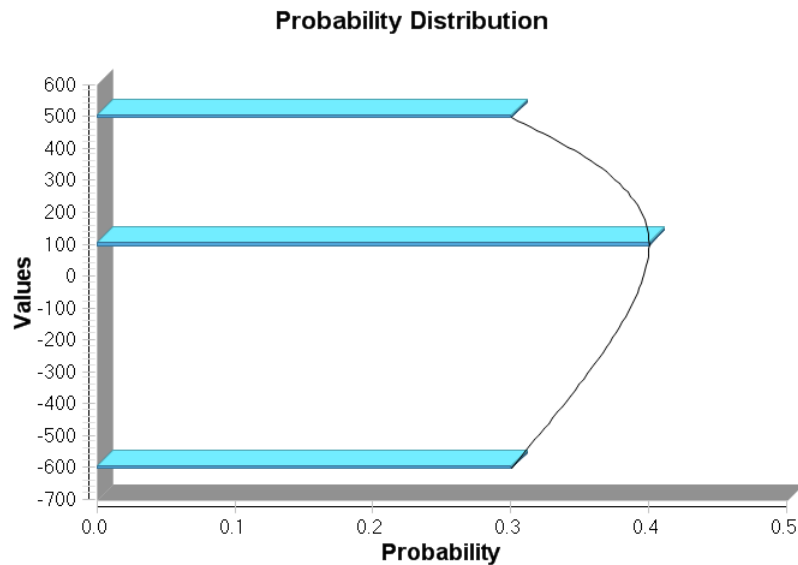
The first tab is labeled "Select Chart Type". This allows you to change the chart type from "Bar" to some other format. In the case of a Probability Distribution, you probably would not change the chart type, but the option is there.

Within the selected chart type, there are subtypes and other properties. For example, you could change the Dimension option to 2D With Depth, and the chart would look like this.



Bar graph with depth

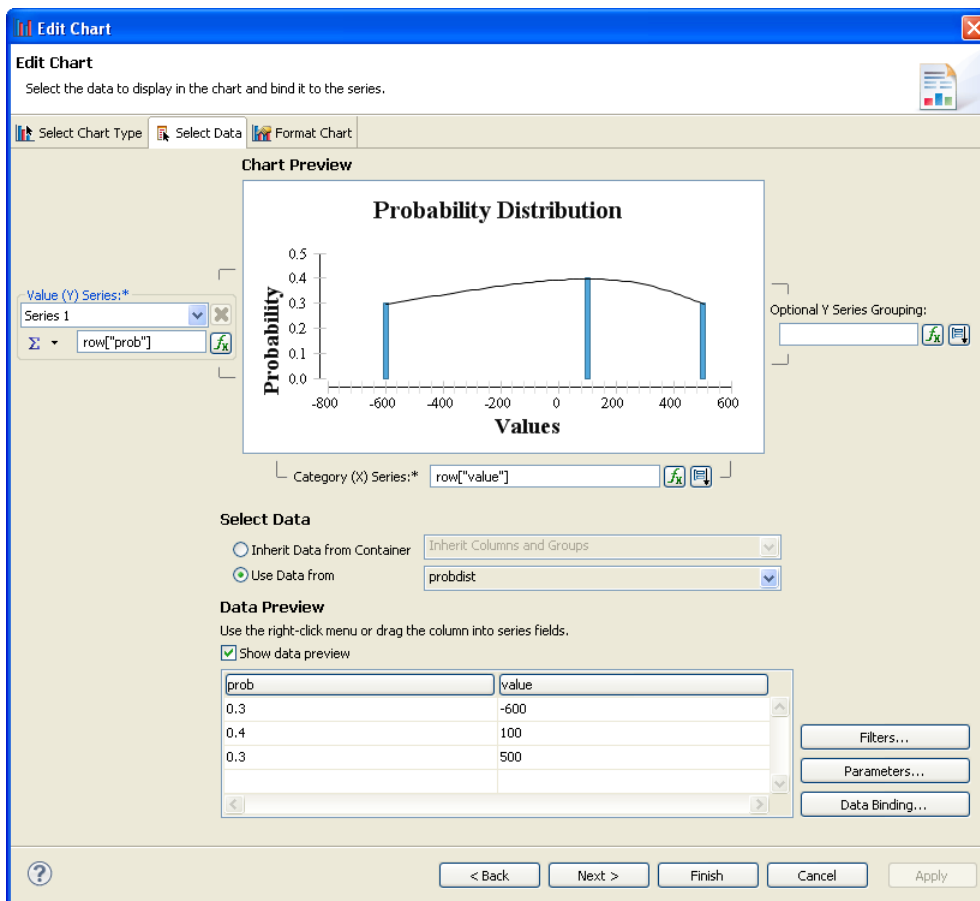
You could also change the orientation by flipping the axes. Then the graph would look like this.



Bar graph with flipped axes

9.2.2 Changing the chart data

The second tab in the Edit Chart dialog is "Select Data".

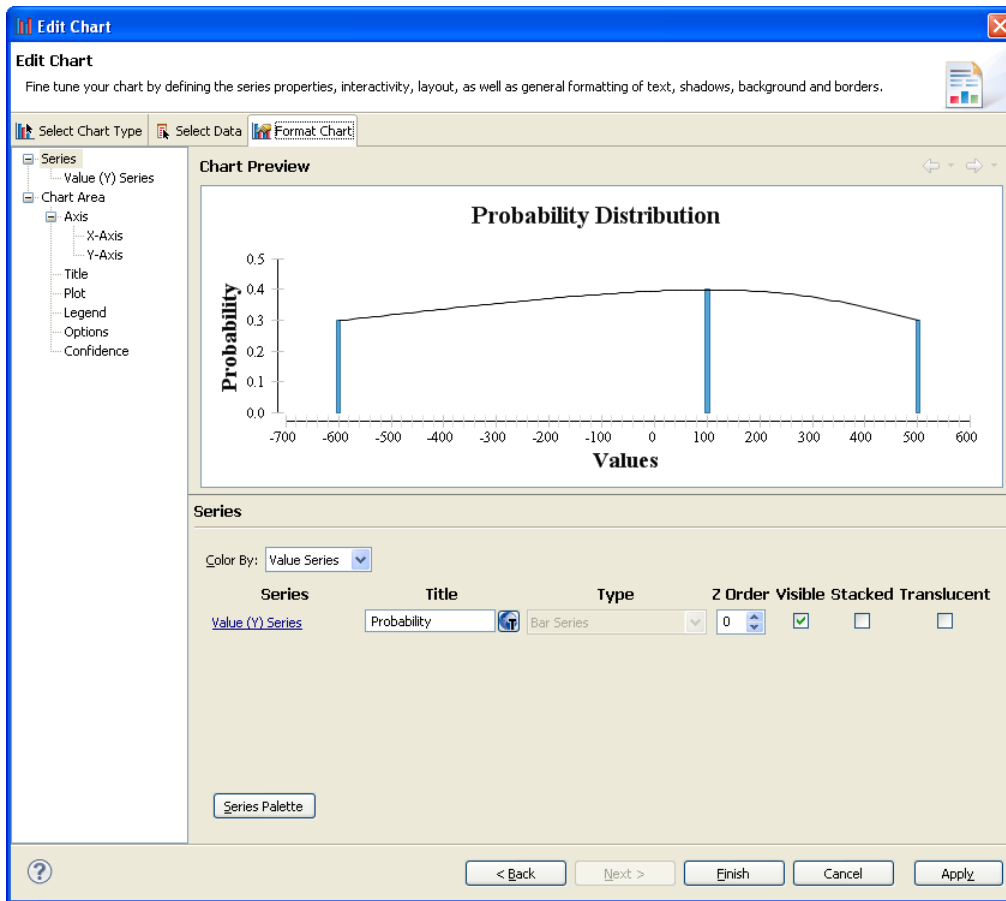


Edit Chart Dialog - Select Data tab

These options allow you to select different data from the graph's text report. Note that the text report data is presented in this tab when the "Show data preview" option is selected. In general, TreeAge Pro will select the data that is appropriate for the graph. These options should only be used with care.

9.2.3 Changing the chart format

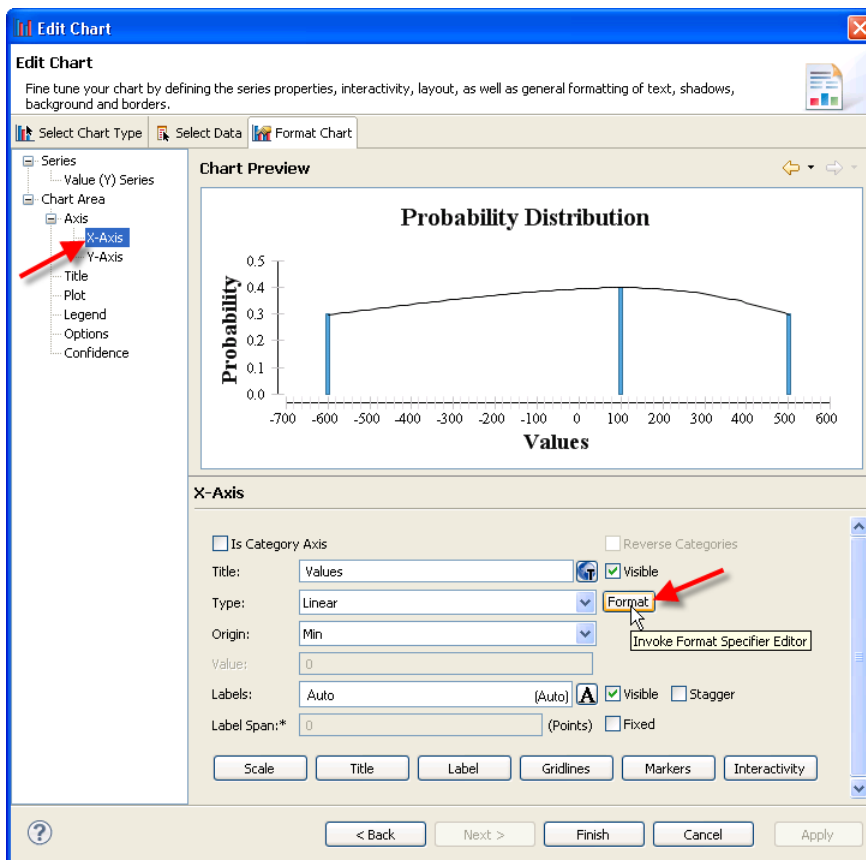
The third tab in the Edit Chart dialog is "Format Chart".



Edit Chart Dialog - Format Chart tab

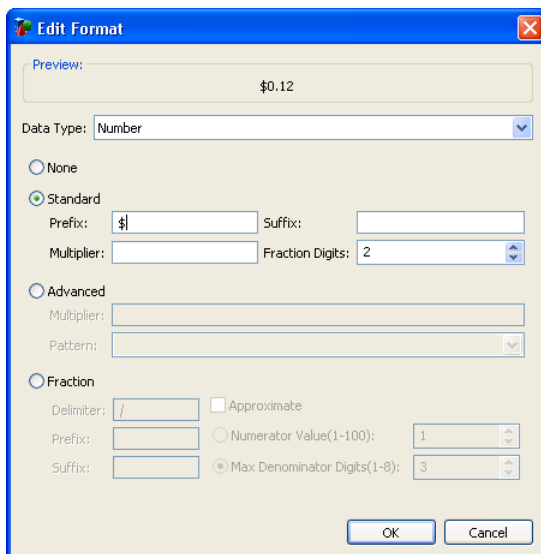
The Format Chart tab allows you to change chart attributes like the chart title, hide/show legend, etc via the selections in the left frame.

In addition, you can change formatting options related to the X-axis and Y-axis by selecting the axis in the left frame, then clicking the Format button.



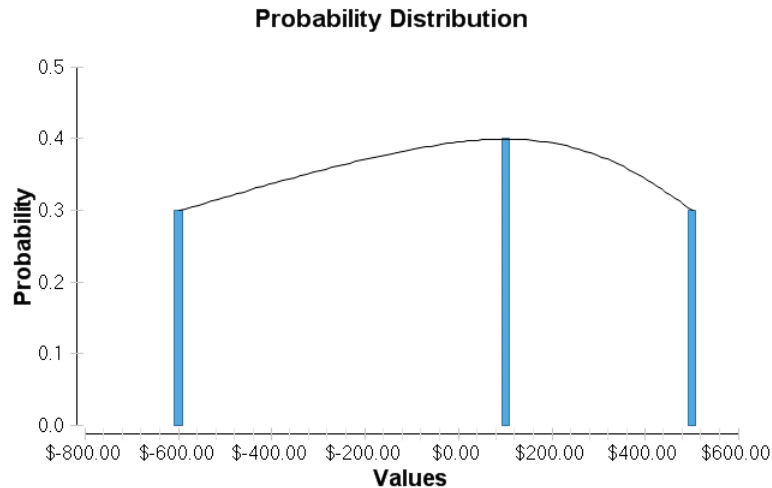
Edit Chart Dialog - Format axis

That will open the Edit Format dialog, which allows you to edit the formatting of the selected axis.



Edit Format Dialog

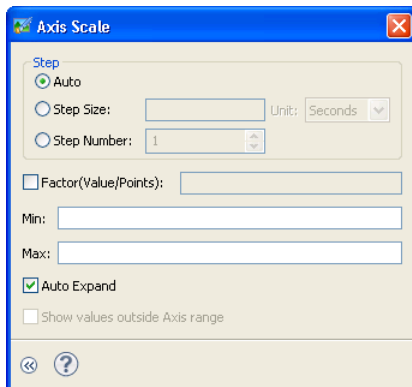
After the changes are applied, the graph's X-axis is reformatted.



Bar graph with formatted X-axis

9.2.4 Changing the chart data scale

You can also change the data scale by clicking the Scale button with one of the axes selected. This allows you to change the minimum, maximum and step values for the selected axis.

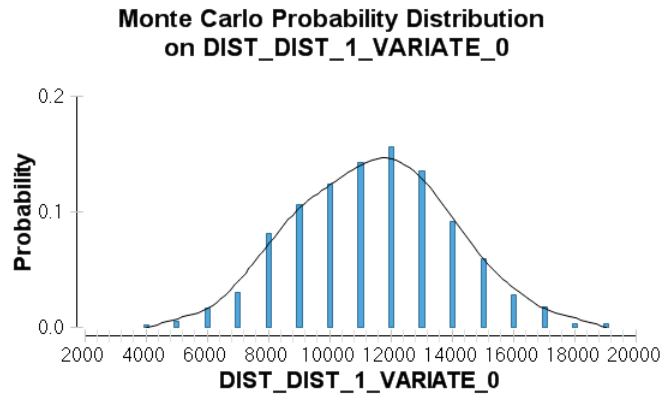


Axis Scale Dialog

9.3 Customizing probability distribution graphs

Although a probability distribution was used to illustrate general graph customization options, there are also some options specific to probability distributions. Specifically, the number of bars in the probability distribution can be adjusted.

The following graph was generated from a Monte Carlo simulation. Specifically, this graph shows the distribution of samples from a model's input distribution.

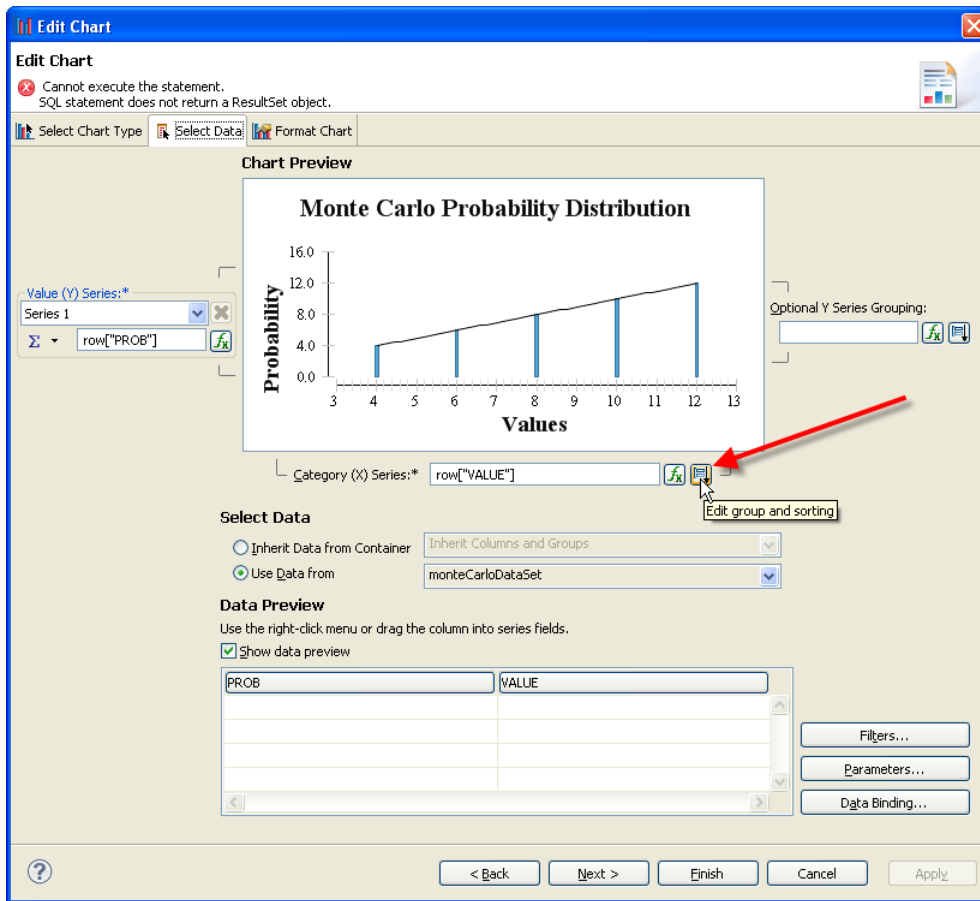


Probability Distribution - Sampling distribution graph

In TreeAge Pro, most probability distributions will include a Redo Histogram link to the right of the graph. This allows you to redraw the graph with more or fewer bars. You can also control the number of bars more precisely through the Edit Chart link as described below.

To show fewer bars:

- Click the Edit Graph link to the right of the graph.
- Click on the Select Data tab in the Edit Chart dialog.
- Click on the Edit group and sorting icon next to the x-axis (see below).
- In the grouping area of the Group and sorting dialog, check the Enabled button and enter an interval value.



Edit Chart Dialog - Group and sorting icon

Group and sorting

Data Sorting: Ascending

Sort On: row["VALUE"]

Grouping

☒ Enabled

Type: Numeric

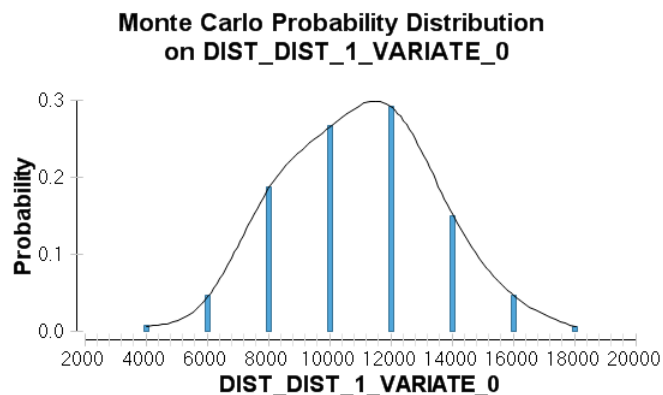
Interval: 2000

Aggregate Expression: Sum

OK Cancel

Group and sorting dialog

The original graph showed a bar for every 1000. The entry above changed this interval to 2000. Note that the customized graph now has fewer bars with higher probabilities.

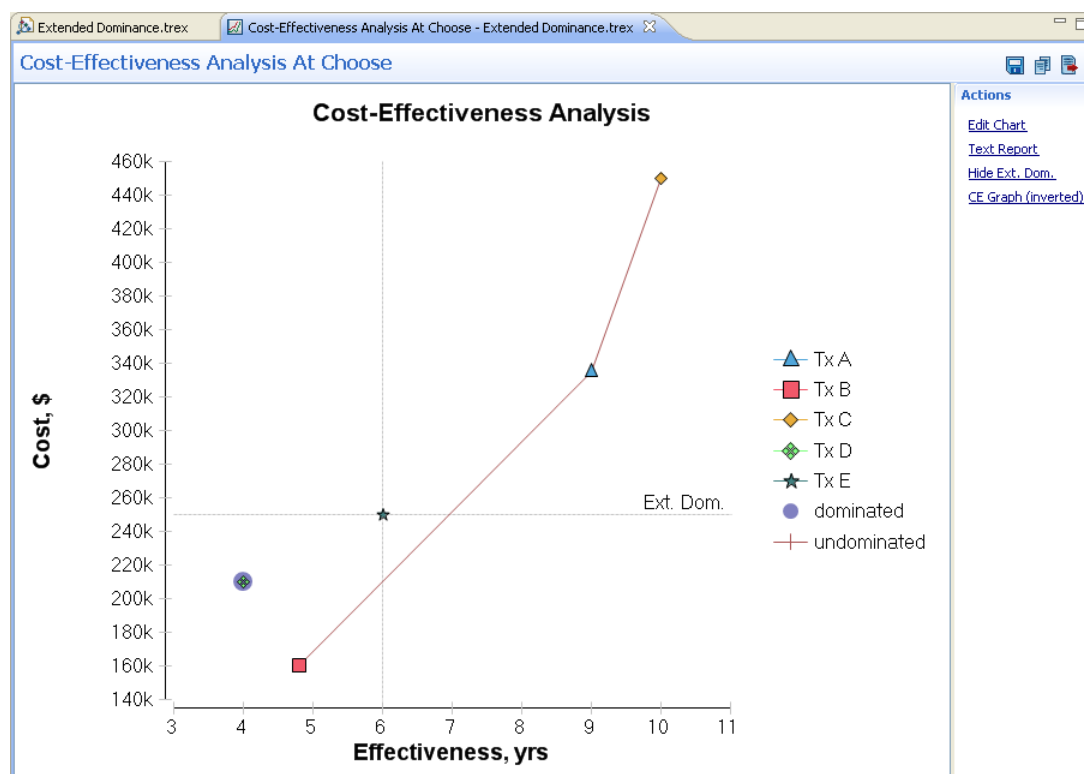


Probability Distribution - Sampling distribution graph - customized intervals

9.4 Customizing cost-effectiveness graphs

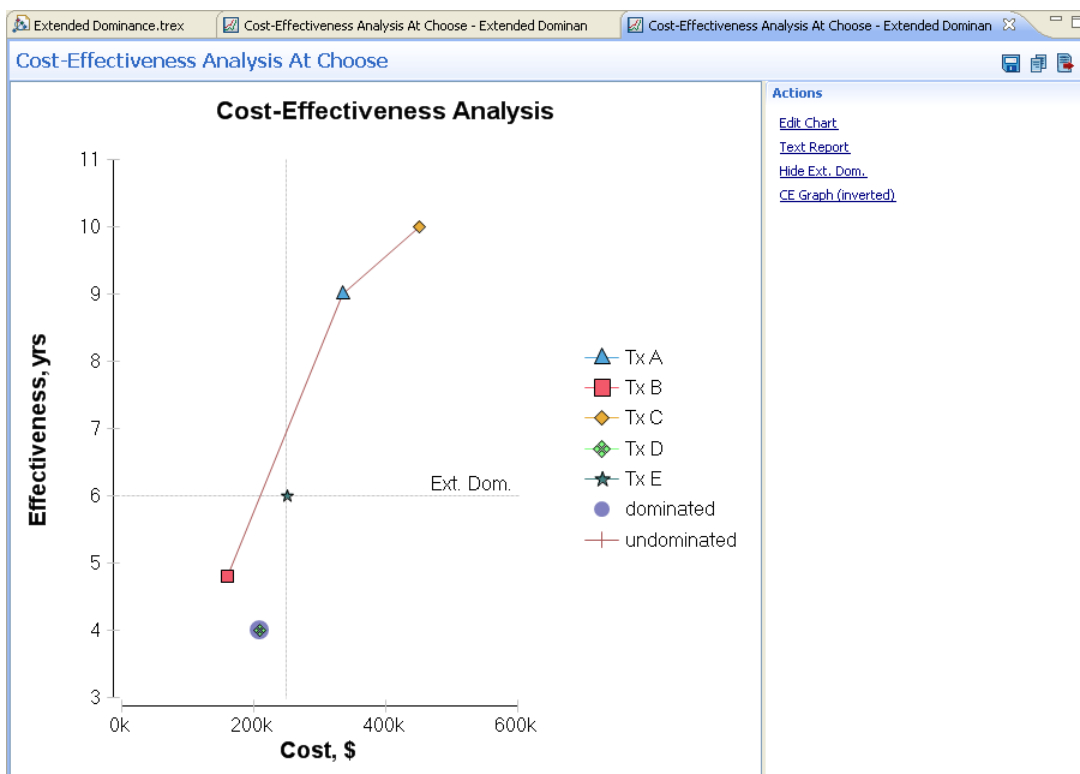
This section describes customization options specific to cost-effectiveness graphs.

The graph below was generated by running cost-effectiveness analysis from the root node of the Health Care tutorial example tree "Extended Dominance".



Cost-Effectiveness Graph

Even before the graph is generated, you are offered the option to show the graph with options inverted. You can also generate the inverted graph by clicking the "CE Graph (inverted)" link. The inverted graph is presented below.



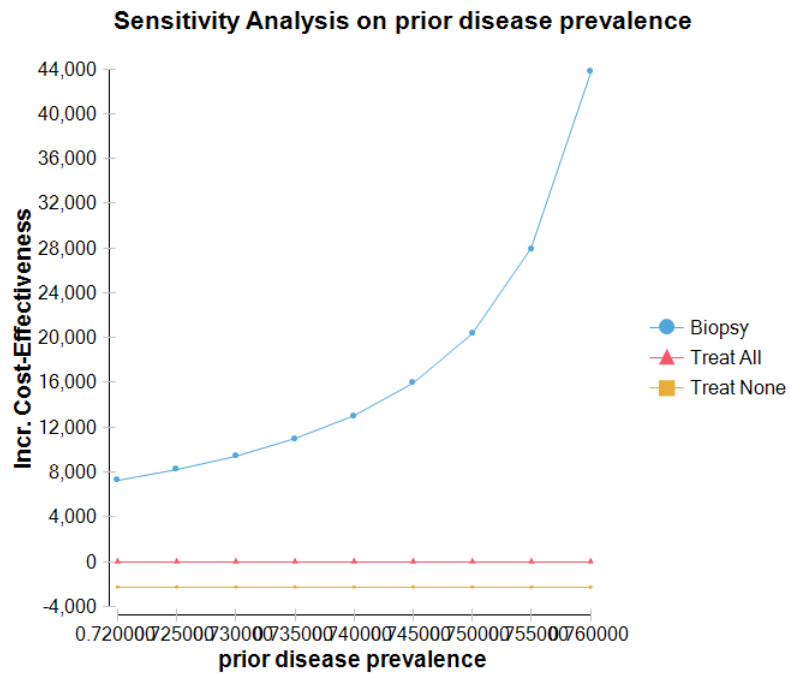
Cost-Effectiveness Graph - Inverted

Another option on this graph is to hide extended dominance lines. Clicking on that link removes the horizontal and vertical lines associated with extended dominance.

Cost-effectiveness graphs show not only points for each strategy, but also use markers for dominated strategies versus undominated strategies. Because of this, changing the chart type and other formatting options can cause unexpected effects on the graph. You can still make changes to scale and most other graph properties.

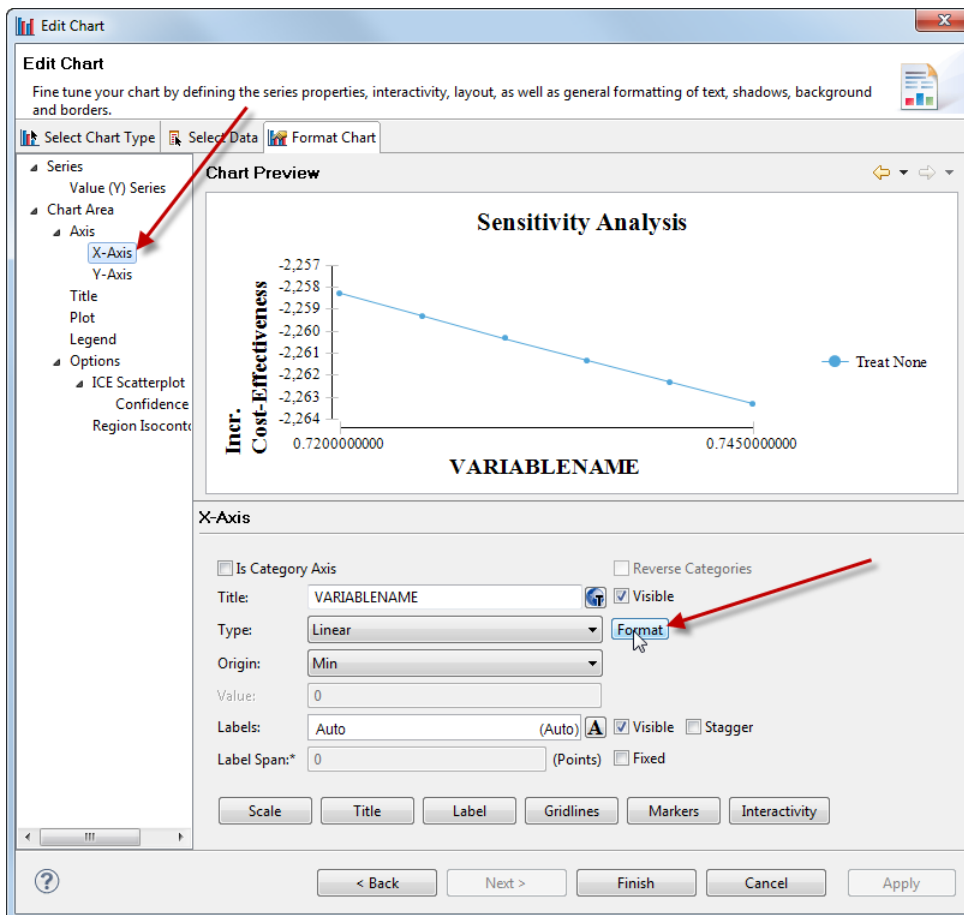
9.5 Customizing line graphs

This section describes customization options specific to line graphs. The following graph was generated from a CE one-way sensitivity analysis. Specifically, this is the x vs ICER graph.



Line graph

Note that the x-axis labels are impossible to read because the values are formatted with too many decimal places. This can be adjusted via the Edit Chart Dialog, Format Chart Tab.



Edit Chart Dialog - X-axis formatting

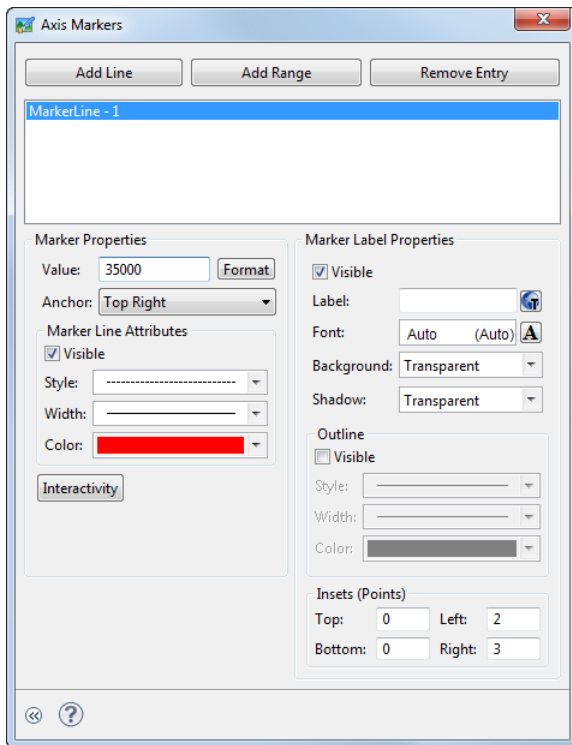
Select the X-Axis in the left pane then click the Format button. Then change the Fraction Digits to 2. See below.

The screenshot shows the 'Edit Format' dialog box. The 'Preview' section shows the value '0.12'. The 'Data Type' is set to 'Number'. The 'Standard' radio button is selected. The 'Prefix' and 'Suffix' fields are empty. The 'Multiplier' field is empty. The 'Fraction Digits' field is set to '2'. The 'Advanced' radio button is unselected. The 'Fraction' radio button is selected. The 'Delimiter' is set to '/'. The 'Approximate' checkbox is unchecked. The 'Prefix' and 'Suffix' fields are empty. The 'Numerator Value(1-100)' field is set to '1'. The 'Max Denominator Digits(1-8)' field is set to '3'. The 'OK' and 'Cancel' buttons are at the bottom.

Edit Chart Dialog - X-axis - Edit Format dialog

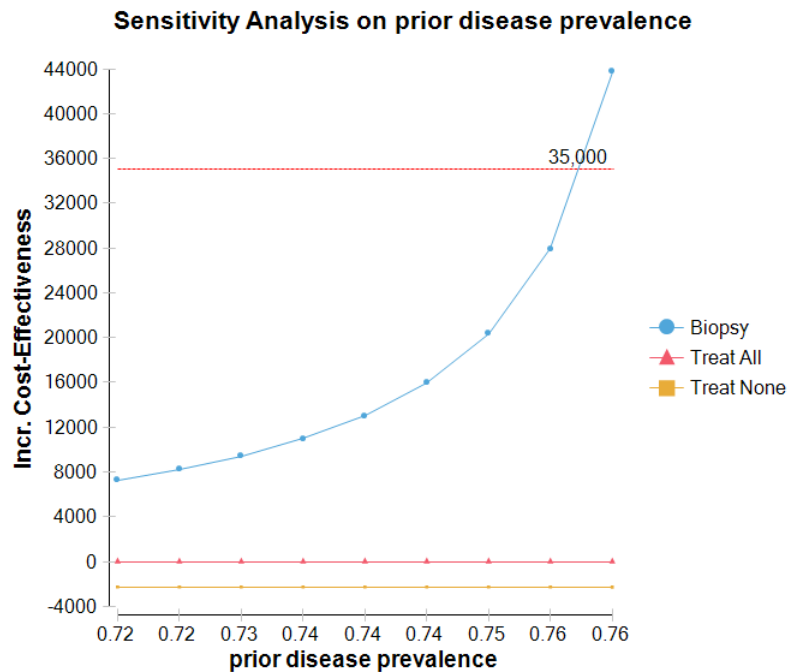
You could also use this dialog to add a custom prefix or suffix to the values.

Let's say you also wanted to add a horizontal line to the graph to highlight the ICER value of 35,000. Select the X-Axis in the left pane then click the Markers button. In the Axis Markers dialog, click the Add Line button, then set the value and properties of the line.



Edit Chart Dialog - X-axis - Axis Markers dialog

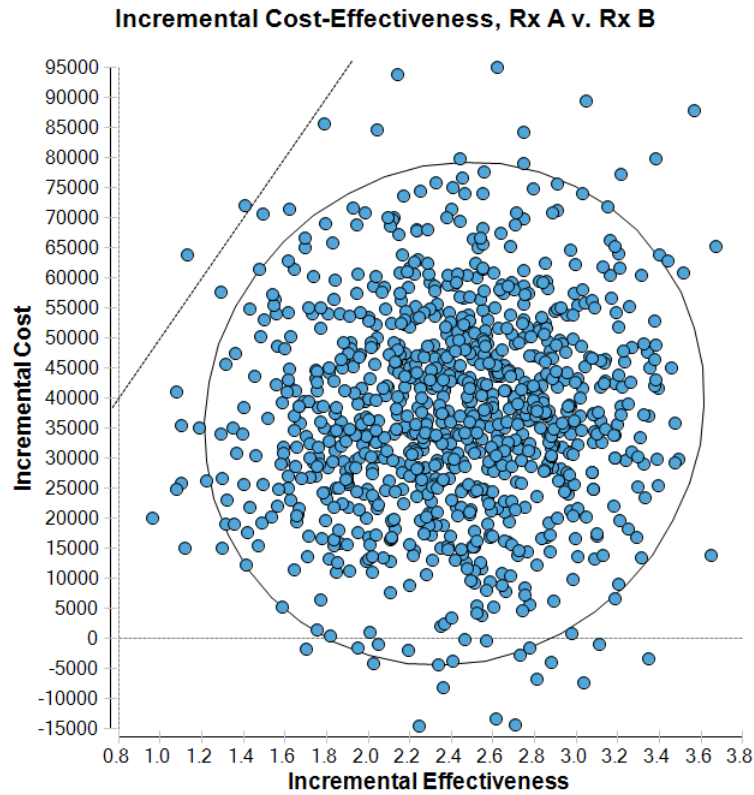
The example above creates a red horizontal line at ICER value 35,000. The combination of the changes above generates the following graph.



Line graph - modified

9.6 Customizing scatterplot graphs

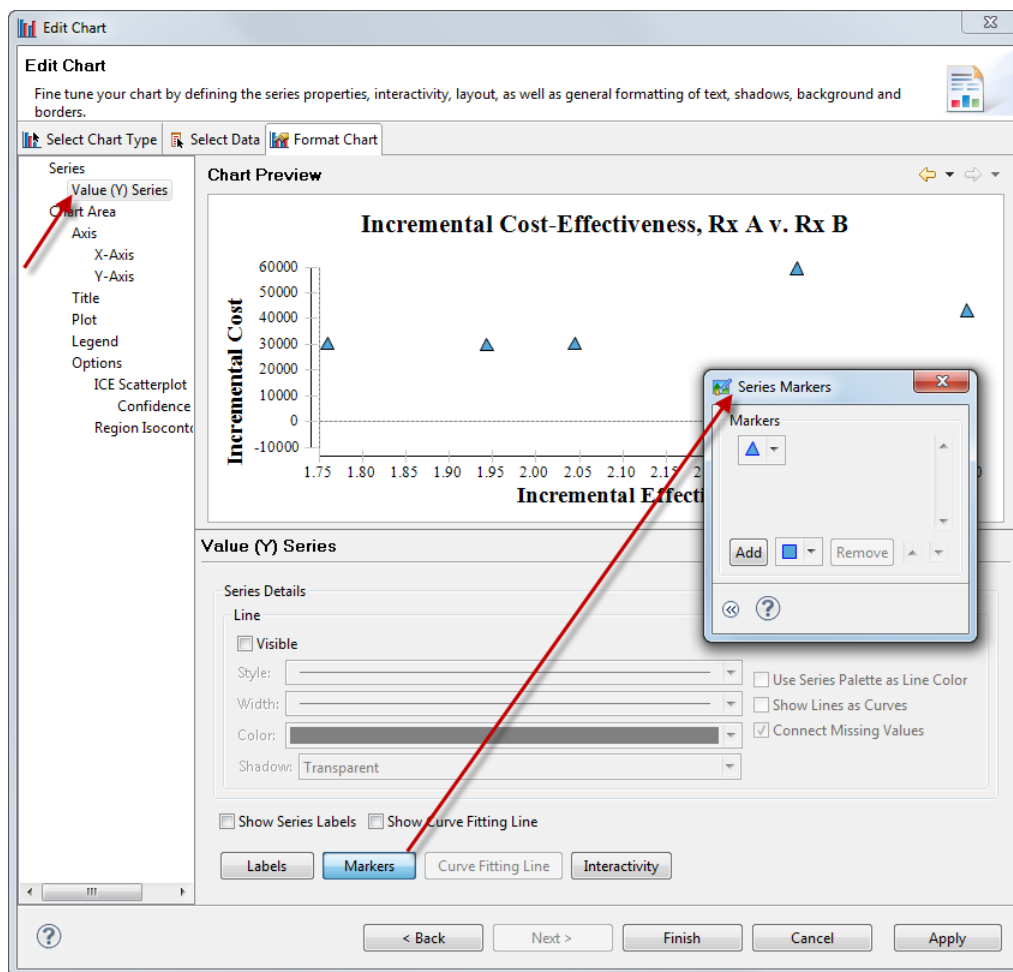
This section describes customization options specific to scatterplot graphs. The graph below was generated from a probabilistic sensitivity analysis simulation on the Health Care tutorial example model CE Markov Sampling. Specifically, the graph is an ICE scatterplot comparing Rx A to Rx B.



ICE scatterplot

You might want to change the X-axis scale to show the 0 value for Incremental Effectiveness. This is described earlier in this chapter.

You might also want to change the plot marker format in the Format Chart tab by selecting the appropriate series and clicking the Marker button. In the example below, the Series markers are changed from circles to triangles.

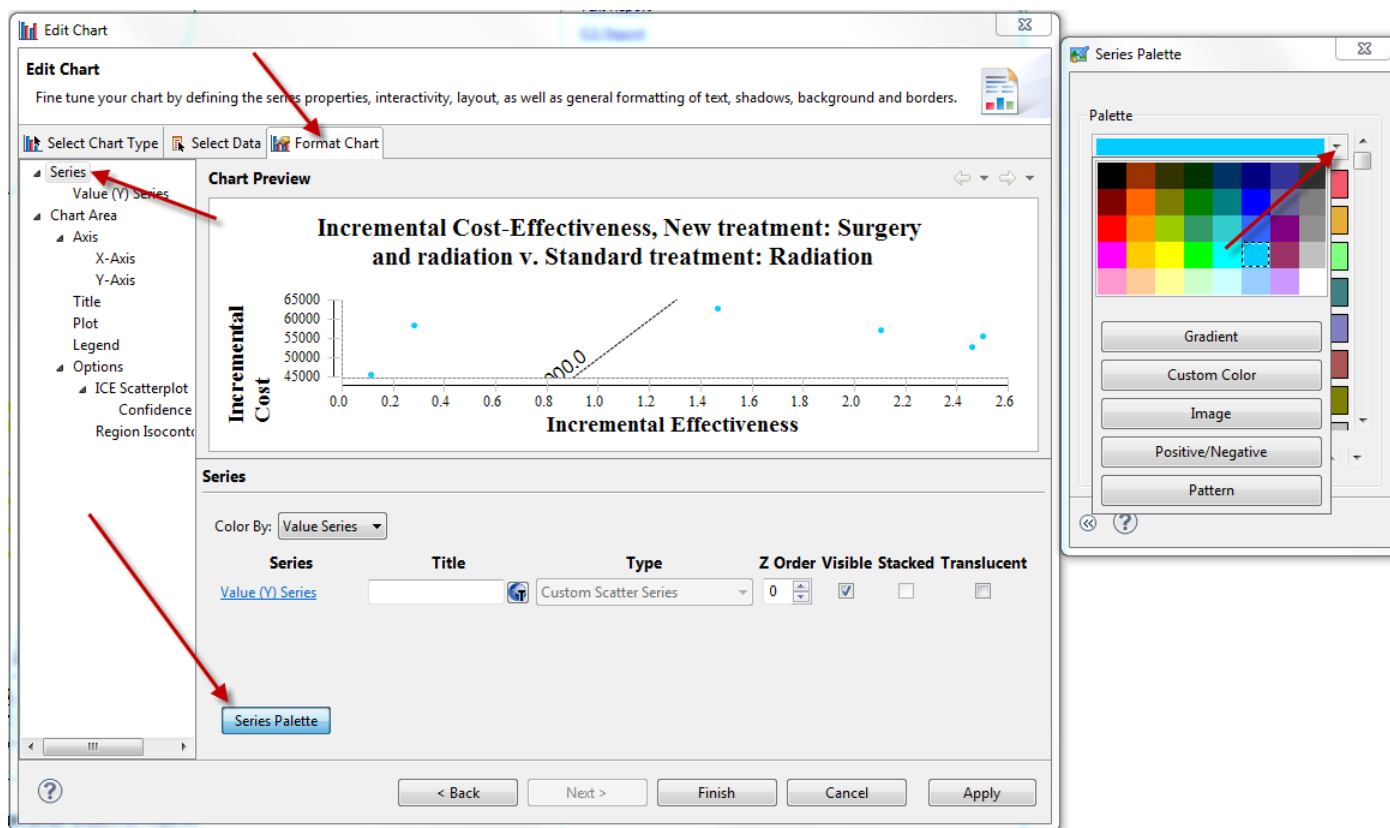


Edit scatterplot - change marker type

Note that if the scatterplot has more than one data series, change the second, third, etc. marker type in the Series Marker dialog to adjust those series.

To change the color of the points in the scatterplot:

- Click the Edit Chart link.
- Select the Format Chart tab.
- Select the category Series (not Value (Y) Series).
- Click the Series Palette button at the bottom of the dialog (may need to scroll down).
- Click on the down arrow to the right of the first palette and select the color you want.
- Click Finish in the main Edit Chart dialog.



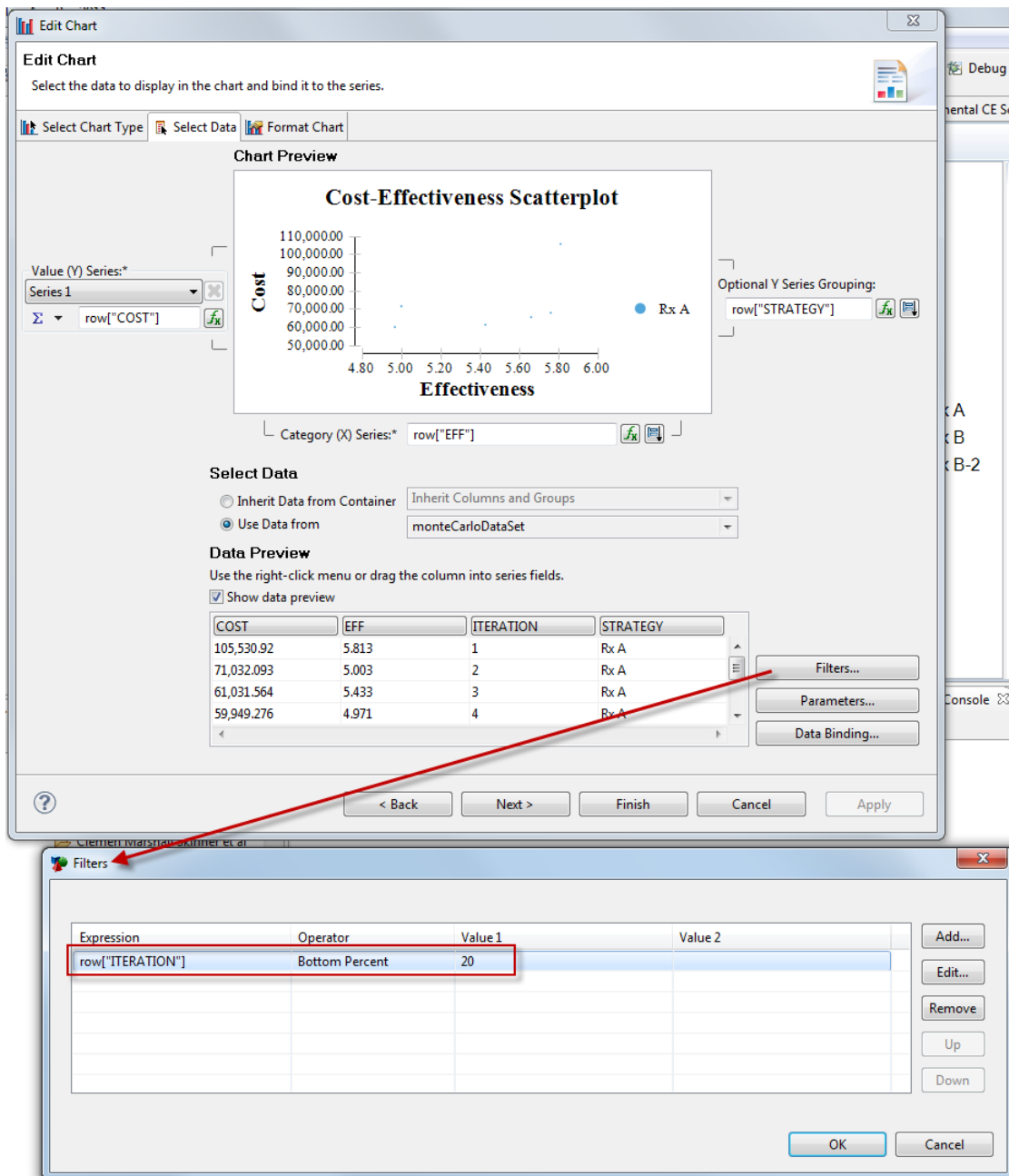
Edit scatterplot - change series color

You might also want to reduce the number of points presented in the scatterplot if the scatter is too dense.

- You can do this by setting a filter on the data points in the Select Data tab.
- Click the Filters button.
- Then add a filter to limit the data points to be displayed.

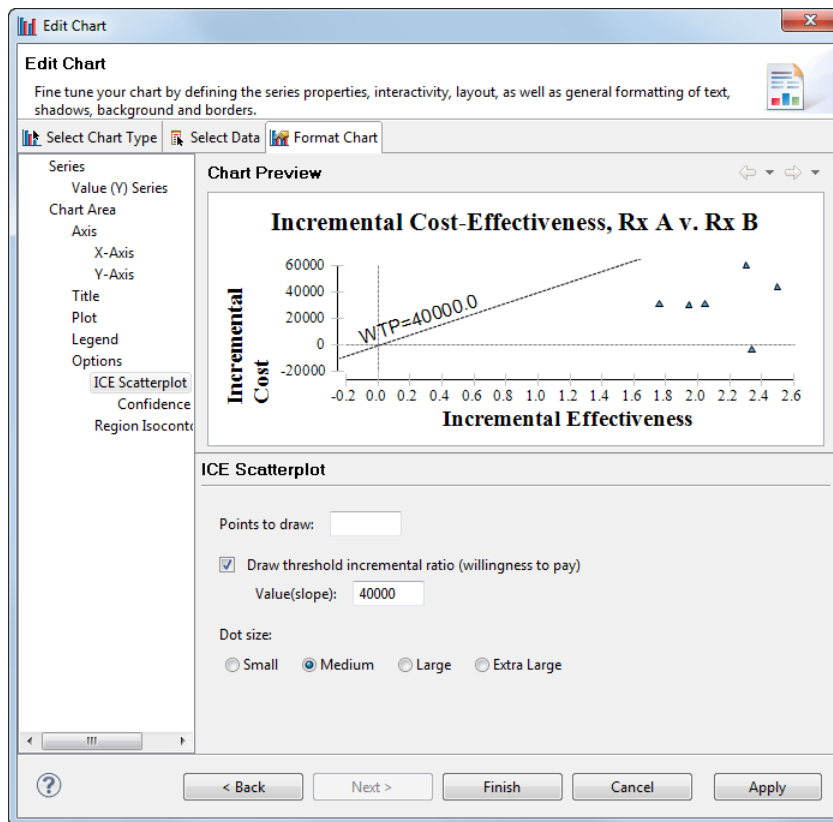
How to reduce the number of points presented:

In the example below, a filter was added to show only the bottom 20 pct of the simulation iterations based on the iteration number. The iteration number is not correlated with any output measure, so you introduce no bias into the new graphical output.



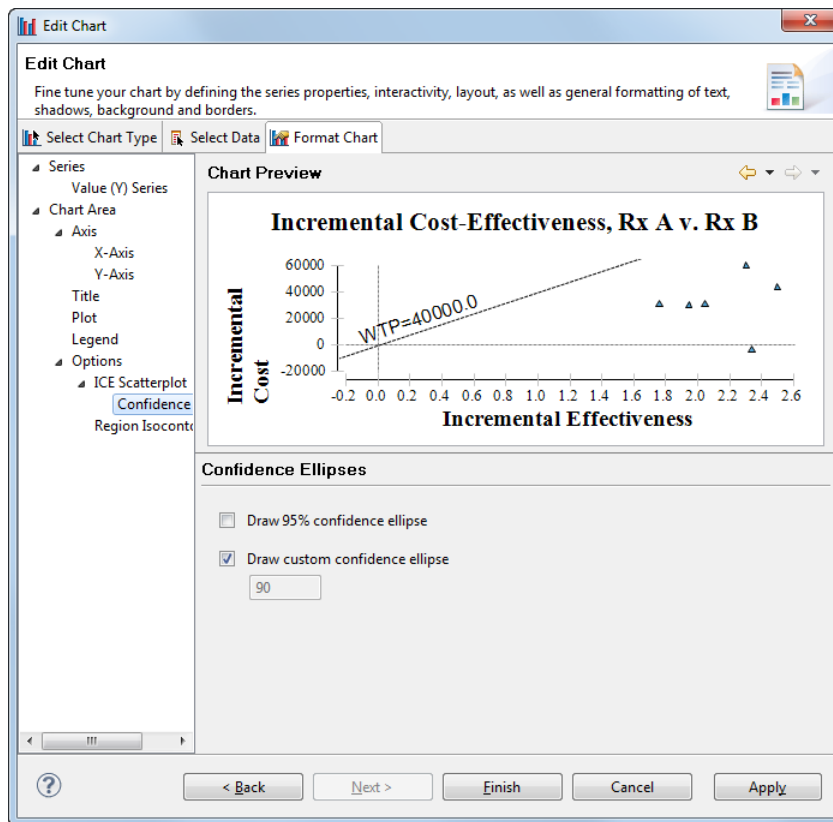
Edit scatterplot - filter data

Beyond standard scatterplot options, the ICE scatterplot has a few specific options that are available under the category ICE Scatterplot in the left frame. This allows you to change the number of points to draw, whether to draw the ICER line, the ICER line's slope and the size of the dots in the scatterplot. See below.



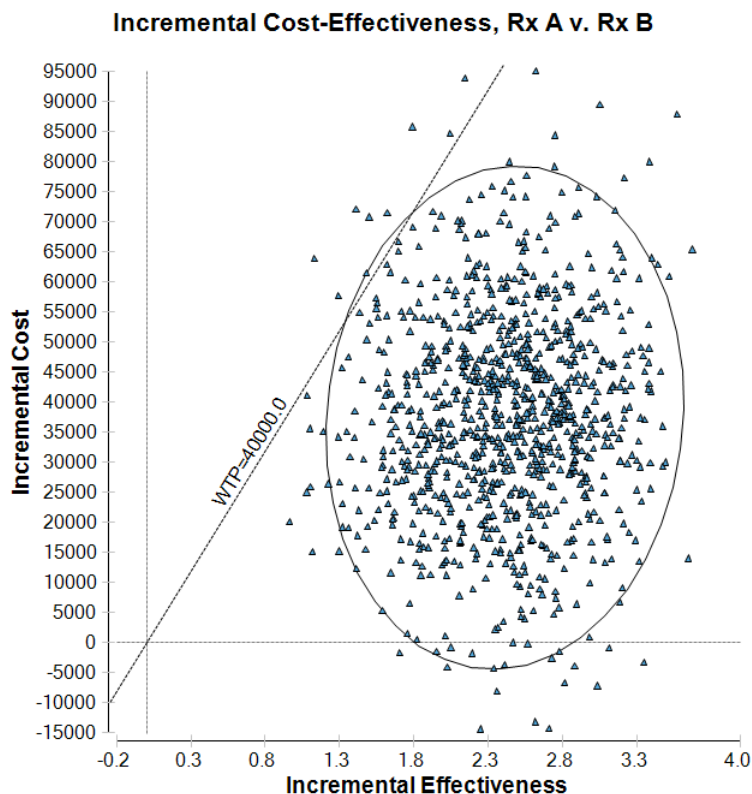
Edit scatterplot - options specific to the ICE scatterplot

The Confidence Interval subcategory allows you to create a custom confidence interval rather than using the default 95% confidence interval. See below.



Edit scatterplot - confidence interval

The net result of all these changes yields the following graph format.



ICE scatterplot - customized

9.7 Customizing tornado diagrams

Note that to change axis information like Scale, Formatting, etc., the horizontal access is modified through the Y-axis settings.

For example, to change the scale of the horizontal access. Choose the category Chart Area > Axis > Y-Axis, then click the Scale button.

9.8 Customizing region graphs

To change the pattern/color of the regions in a region graph:

- Click the Edit chart link.
- Select the Format chart tab.
- Choose the category "Series".
- Click the Series Palette button.
- Click on the first bar and an option arrow will appear at the right of the bar.
- Click on this arrow to change first bar to the desired color or pattern.
- Repeat the prior two steps for additional entries in the palette list as needed.

9.9 Printing, exporting and saving graphs

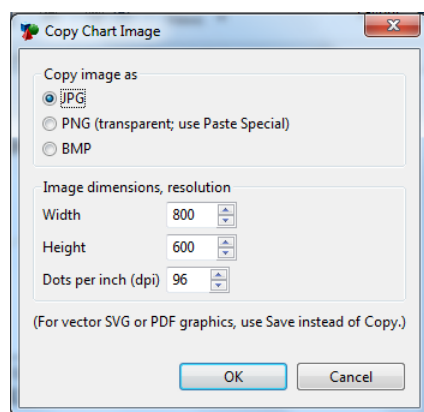
Graphs can be printed, exported and saved via the TreeAge Pro menu when the graph is selected.

To print a graph:


- Select the graph.
- Choose File > Print from the menu.

To copy a graph to insert directly into a document:

- Select the graph.
- To copy the graph and paste a copy into a document, choose from the menu Edit > Copy OR click the Copy icon to the top-right of the graph pane.
- The Copy Chart Image appears, as in the figure below, to select the type of image you want to copy as (jpeg, png, bmp) and to select the resolution.

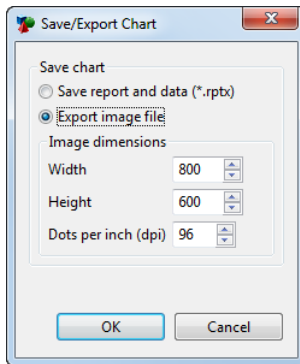
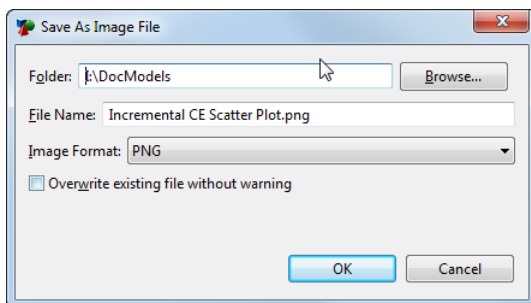


Copy Chart Image appears when copying a graph to insert directly into a document.

 For a sharper image, change width, height and DPI to 1600, 1200 and 300, respectively.

To create an image file from a graph:

- Select the graph.
- Choose File > Save from the menu.
- Choose "Export image file", and enter the appropriate image options in the Save/Export Chart Dialog. (See below)
- Enter the file name, location and format in the Save As Image Dialog. (See below)

**Save/Export Chart Dialog****Save As Image File Dialog**

To save a graph in its native TreeAge Pro format:

- Select the graph.
- Choose File > Save from the menu.
- Choose "Save report and data" in the Save/Export Chart Dialog. (See above)
- Select the file name and location.

10. Tree Calculation Methods and Preferences

Any single tree may be calculated and evaluated in a variety of ways, simply by switching between different payoffs, or from single- to multi-attribute calculations. This chapter covers tree calculation method preferences, including two kinds of multi-attribute calculation methods.

An additional multi-attribute calculation method, Cost-Effectiveness, is available to users of the Healthcare module; refer to the Building and Analyzing Cost-Effectiveness Models for details.

Other chapters cover the other preferences categories; refer to the Preferences Chapter for an overview of all preferences.

10.1 Changing what the tree calculates

Some basic tree calculation preferences were introduced at the end of the Decision Tree Tutorial Chapter and the beginning of the Analyzing Decision Trees Chapter:

- the calculation method
- the active payoff(s)
- the optimal path criterion
- numeric formatting

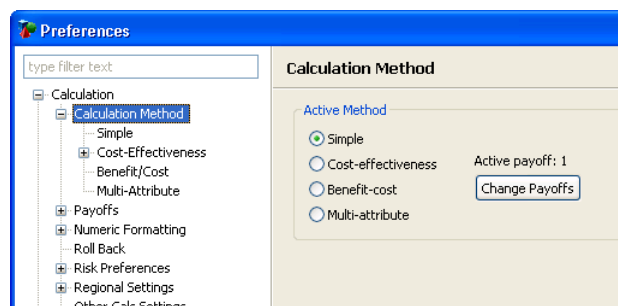


Note that a set of Tree Preference Values can be stored in a Preference Set.

To view in detail the tree's current calculation method, or to make changes to these settings, open the Preferences dialog.

To view/edit the calculation method preferences:

- Choose Tree > Tree Preferences from the menu.
- Select the category Calculation > Calculation Method.



Tree Preferences - Calculation Method

10.1.1 Selecting calculation method

The choice of calculation method determines the formula used to calculate values for nodes in your tree. There are two kinds of calculation methods:

- Simple, single-attribute calculations;
- Multiple-attribute calculations, including: Benefit-Cost and Multi-Attribute (weighted);
- Healthcare module users can use a third form of multi-attribute calculations, Cost-Effectiveness.

Simple calculations are just that — expected values are calculated for the nodes in the tree based simply on the values in the active payoff set. If, as described above, your tree included multiple attributes — such as monetary benefits in payoff #1 and costs in payoff #2 — the two sets of payoff values could be combined in a single calculation using the Benefit-Cost calculation method's formula.

To change the calculation method:

- Roll back must be turned off.
- With the Preferences dialog open to the Calculation Method category, select the radio button for the appropriate calculation method.
- If you select a calculation method for the first time in a particular tree, the settings for payoffs, optimal path criterion, and numeric format may need to be modified.



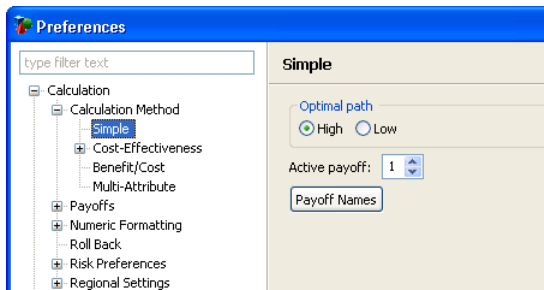
Calculation method notes:

- Changing calculation method (or other) preferences will not affect the content of your tree. Values or formulas already entered in payoff #1 will not be lost by switching the active payoff set from #1 to #2, for example. Only the calculation and display of values will be affected; the changes to the calculation preferences can be reversed at any time.
- When the Preferences dialog is closed, TreeAge Pro updates the information in the status bar. If you have switched Simple calculations to use payoff 3, "Payoff 3" will appear in the status bar. If you select Benefit-Cost using payoffs 4 and 2, the status bar will read "B-C, 4-2." If you select Multi-Attribute, then "MultiAttr" will show in the status bar.

Additional Tree Preference categories contained within the Calculation Method group are used to specify preferences associated with each calculation method. These categories are described in subsequent sections.

10.1.2 Calculation method "Simple"

The Simple Tree Preferences category is presented below.



Tree Preferences - Simple Calculation Method

Optimal path

All calculation methods in TreeAge Pro require you to specify how decisions should be made in the tree (i.e., during roll back). This is done using the option labeled *Optimal path*.


The default optimal path criterion used for new trees is High, unless you save new default Preference Set. This setting would be changed if, for example, you were modeling project costs (to find the strategy with the lowest expected cost).

To set the optimal path criterion:

- Select the tree and press the F11 key to open the Tree Preferences dialog.
- Select the category Simple within the Calculation Method group.
- Next to the label Optimal path, select the radio button for either High (maximize expected values) or Low (minimize expected values).


For a tree set to High (e.g., one whose payoffs are in terms of life expectancy), at each decision node TreeAge Pro will select the alternative with the *highest* numeric value.

For a tree set to Low (e.g., one whose payoff formula is in terms of costs), the alternative with the *lowest* numeric value is selected.

 Each of the nine payoff sets stores its own optimal path criterion for Simple calculations, and each form of multi-attribute calculations also has its own optimal path setting.

Active payoff

In TreeAge Pro, you can enter any number of payoffs at each terminal node. In a Simple analysis, you can use any one of the enabled payoff/reward sets. The *Active payoff* tree preference is used to select the active payoff for the Simple Calculation Method.

 The *Active payoff* option will allow selection of any enabled payoff set. Based on the default Preference Set, two payoff sets are enabled, but this can be changed in the Payoffs Tree Preferences category.

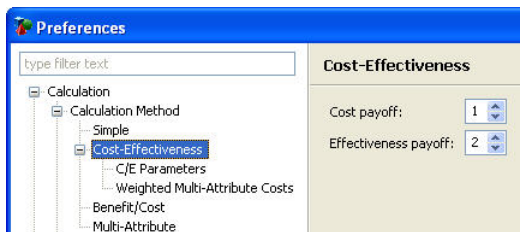
To change the active payoff:

- In the Calculation Method/Simple preferences, select a payoff set in the Active payoff field.

- When you change to a new Active payoff for the first time in a tree, ensure that you select the appropriate optimal path criterion and numeric formatting for that payoff.

10.1.3 Calculation method "Cost-Effectiveness"

The Cost-Effectiveness Tree Preferences category is presented below. Note that this calculation method is only available if your license includes the Healthcare Module.



Tree Preferences - Calculation Method Cost-Effectiveness

Cost payoff

Select the payoff set to use for cost calculations from among the enabled payoff sets.

Effectiveness payoff

Select the payoff set to use for effectiveness calculations from among the enabled payoff sets.

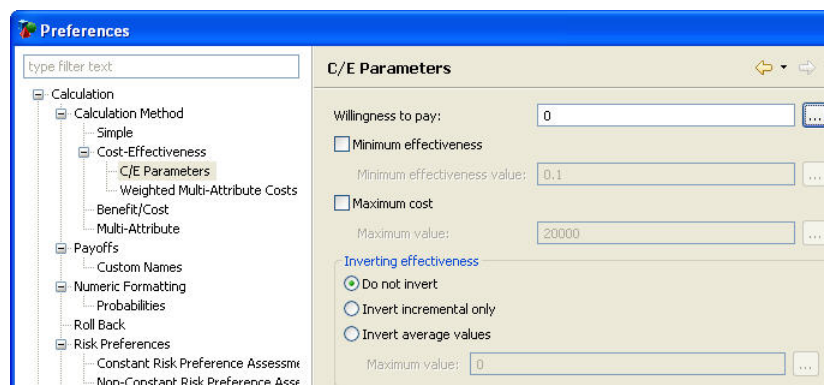
Both the above controls work the same as the Active payoff in the Calculation Method/Simple preferences.

Additional cost-effectiveness options

Additional Tree Preferences related to the cost-effectiveness calculation method are described in the next two sections.

10.1.4 Calculation method "Cost-Effectiveness", C/E Parameters

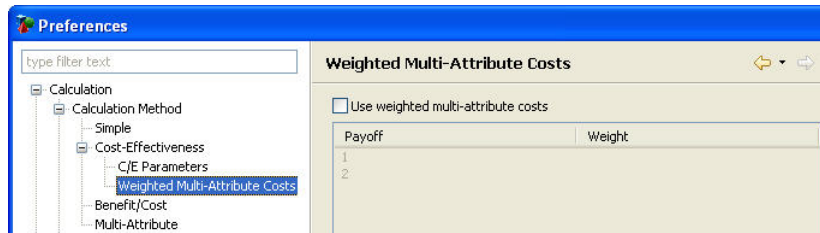
The Cost-Effectiveness, C/E Parameters Tree Preferences category is presented below.



Tree Preferences - C/E Parameters

10.1.5 Calculation method "Cost Effectiveness", Weighted Multi-Attribute

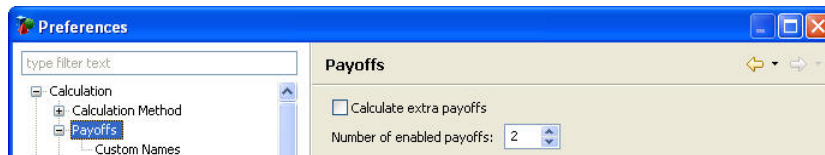
Content of the new section ...



Tree Preferences - Weighted Multi-Attribute Costs

10.1.6 Enabling payoffs

TreeAge Pro supports any number of distinct payoffs sets (attributes) in a tree. By default, only two payoff sets are enabled, but this can be changed via the Payoffs Tree Preferences category.



Tree Preferences - Payoffs

Calculate extra payoffs

Check this box to calculate all enabled payoffs rather than calculating only the active payoffs for the Calculation Method. This option is described in detail in the next section.

Number of enabled payoffs

Select the number of payoff sets to enable within the model. For example, if you change the number of enabled payoffs to 5, then payoff sets 1 through 5 will be enabled.

10.1.7 Calculate extra payoffs

It is possible to calculate the tree and optimize at decision nodes based on the current active payoff method and set(s), but report expected values for other payoff sets. For example, you could roll back the tree based on minimizing costs in payoff #1, but also report expected values for some types of utility attributes which you have entered in payoffs #2 and up.

In the Payoffs Tree Preferences category, check the box labeled *Calculate extra payoffs*, and adjust the *Enable payoffs* setting to control how many extra payoff sets will be calculated.

To report extra payoffs in roll back, set up terminal columns using a custom calculation like "Node(N)" where N is the extra payoff. The Node() function can be used in a variety of ways to retrieve expected values.

If *Calculate extra payoffs* is checked, the Monte Carlo simulations and Markov Cohort Analysis (Quick) automatically include the calculated extra payoffs using additional text report columns and distribution graphs. The detailed Markov analysis text report will also include the extra reward sets.



If you activate the calculation of extra payoffs, you must fill in the enabled payoff sets at all terminal nodes, even though they are not specifically required by the current calculation method.

Under Multi-attribute calculations, weight values must be specified for every enabled extra payoff. Use a 0 weight to exclude an extra attribute from the regular, weighted expected value calculation (this will not zero the extra payoff's reported value).

10.2 Calculations using multiple attributes

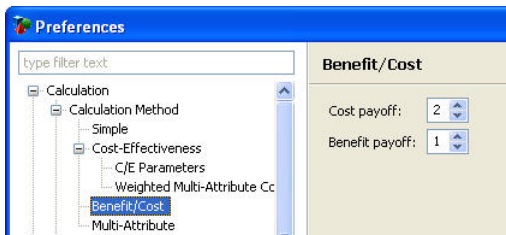
While many analyses will require only one attribute (e.g., cost), some models may have multiple attributes (e.g., benefits and costs) or different perspectives on the same attribute (e.g., societal, personal, or institutional costs of disease).

TreeAge Pro includes two basic calculation methods that combine multiple payoffs: benefit-cost and weighted multi-attribute calculations. (If you have the optional Healthcare module, a third calculation method using multiple payoffs is available: Cost-Effectiveness. See the Healthcare module documentation for details.)

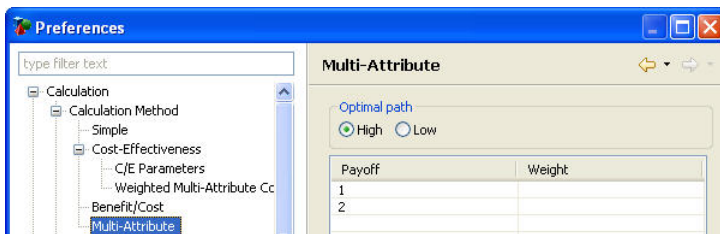
The first step in preparing a multi-attribute model is setting the calculation preferences.

To select a multi-attribute calculation method:

- Select the tree and press the F11 key to open the Tree Preferences dialog.
- Select the category Calculation Method.
- Change the Calculation Method pop-up menu selection from Simple to one of the multi-attribute options.
- If you select Benefit-Cost...
 - Select the Benefit/Cost category (see below).
 - Select two payoff sets to represent the two attributes in your tree.
 - The optimal path will automatically be High (maximize value).
- If you select Multi-Attribute...
 - Select the Multi-Attribute category (see below).
 - Select the Optimal path option.
 - Enter a weight for each enabled payoff.



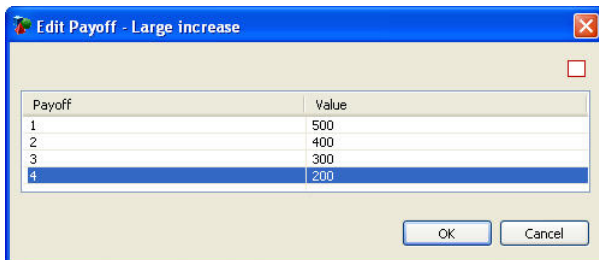
Tree Preferences - Benefit/Cost Calculation Method



Tree Preferences - Multi-Attribute Calculation Method

10.2.1 Entering multiple payoffs

With both kinds of multiple-attribute models — benefit-cost and multi-attribute — you will enter at least two payoffs at each terminal node. The process of entering multiple payoffs values at a terminal node is very simple — the payoff sets in the Enter Payoff window correspond to the Use payoff selections made in the Preferences dialog.



Enter Multiple Payoffs at Terminal Node

If a multi-attribute calculation method is used, the payoff titles in the window will indicate which payoffs are to be used. For benefit-cost, they will be labeled “Benefit” and “Cost.” For weighted multi-attribute models, they will be labeled by payoff set number (as above).

If you require more payoff sets in your model (e.g., for weighted multi-attribute calculations), you can enable additional payoffs.

10.2.2 How benefit-cost calculations are performed

This calculation will subtract the cost of a scenario, represented by one payoff, from its benefit, represented by a different payoff. Therefore, both attributes in a benefit-cost analysis must be measured

in the same monetary units. All analyses available under Simple calculations (1-, 2-, 3-way sensitivity analysis, for example) are also available under the Benefit-Cost calculation method.

This calculation does not divide costs by benefits, as is done in some forms of cost-benefit analyses. To create a custom cost/benefit analysis calculation method, refer to the section on the Calculate extra payoffs preference and the Node() function.

10.2.3 How weighted multi-attribute calculations are performed

Unlike the Benefit-Cost calculation method, the weighted Multi-Attribute calculation method does not require that all attributes/payoffs be assigned using the same units. Instead, a linear equation is set up that combines up to nine payoffs; attributes given a non-zero weighting are made part of the multi-attribute calculations.

For example, if you assigned a weight of 1 to Attribute 1 and a weight of 0.5 to the next three attributes, each terminal node would be evaluated based on the expression $\text{Attribute 1} + 0.5 * \text{Attribute 2} + 0.5 * \text{Attribute 3} + 0.5 * \text{Attribute 4}$.

Variables (see Variables Chapter) can be used in the weighting expressions. This is useful when there is uncertainty concerning how much one factor should be weighted versus another.

Weightings are entered through the Multi-Attribute Tree Preferences category within the Calculation Method group (see earlier section).

All analyses available under Simple calculations are available under Multi-Attribute calculations as well.

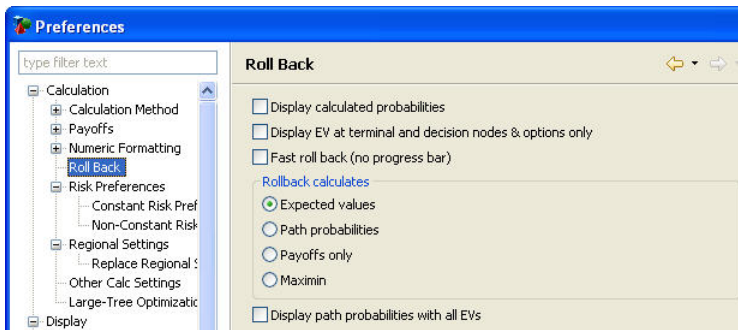
10.3 Roll back analysis options

Before dealing with multiple-attribute analysis methods, let's look at some calculation preferences related to roll back.

Typically, roll back boxes display the expected value of each node, along with the path probability at terminal nodes. TreeAge Pro can display a variety of other calculated values during roll back; these are described below.

To change the quantity to be calculated during rollback:

- Select the tree and press the F11 key to open the Tree Preferences dialog.
- Select the category Calculation Method.
- Select an option from the group labeled *Rollback calculates*.



Tree Preferences - Roll Back

Here is a brief description of the other options:

Expected values: Display the expected values (the default setting) at all nodes.

Payoffs only: Display only the values of terminal nodes. Optimal paths will be indicated using hash marks and colored lines, but no expected values will be displayed.

Path probabilities: Suppress the display of expected values. Path probabilities are calculated for all terminal nodes (does not take into account the optimal path). If there are any decision nodes, the sum of the path probabilities of all terminal nodes will be greater than 1.0.

Maximin: Consider only the most pessimistic possibility at each uncertainty, regardless of probabilities. Then, at each decision point, the best option is selected. More details are provided in the following section.

10.3.1 Maximin and minimax roll back

Maximin calculations may be useful during early stages of certain models, before probabilities have been assigned to the tree. Here, specifically, is how a tree is rolled back under Maximin:

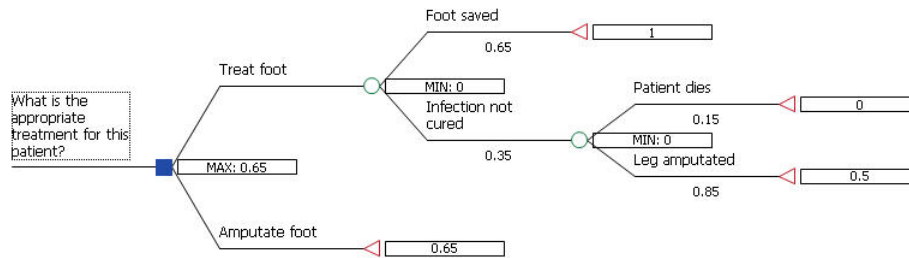
- The value assigned to every chance node is equal to the worst (least optimal) value of any of its potential outcomes. *Probabilities are ignored*, and may be left blank.
- The value assigned to every decision node is equal to that of the best alternative, as usual.



If you use the Options > Reverse Optimal Path command at selected decision nodes, these nodes will work like chance nodes, as described above.

Maximin calculations take a pessimistic view of events. They are based on the idea that one way to deal with risk is to identify the worst case scenario for each alternative and then select the strategy which yields the best of these worst case scenarios.

Below is a picture of the example file Rock Climber using Maximin roll back. The value of the Treat foot strategy is simply its worst outcome, which is the patient dying (utility = 0). The Amputate foot option has no uncertainty, and it is simply equal to its payoff. The decision then maximizes between the available options, as usual.



Rock Climber - Roll back with Maximin

In a Maximin roll back of a tree whose optimal path preference is set to High, the roll back boxes at chance nodes will be labeled MIN: at decision node the boxes will be labeled MAX:, in order to indicate the operation being performed. These prefixes are switched if the optimal path preference is set to low (minimization).

10.4 Regional/international numeric settings

By default, the regional numeric settings specified in your operating system determine which character TreeAge Pro will recognize as a decimal separator when you enter numbers — usually either a “.” or a “,” — and which character, if any, represents a thousands separator.

For example, your computer’s settings are used to decide whether the entered number:

1,375

should be interpreted with *comma = decimal* as

$1 + 375/1000$

or with *comma = thousands separator* as

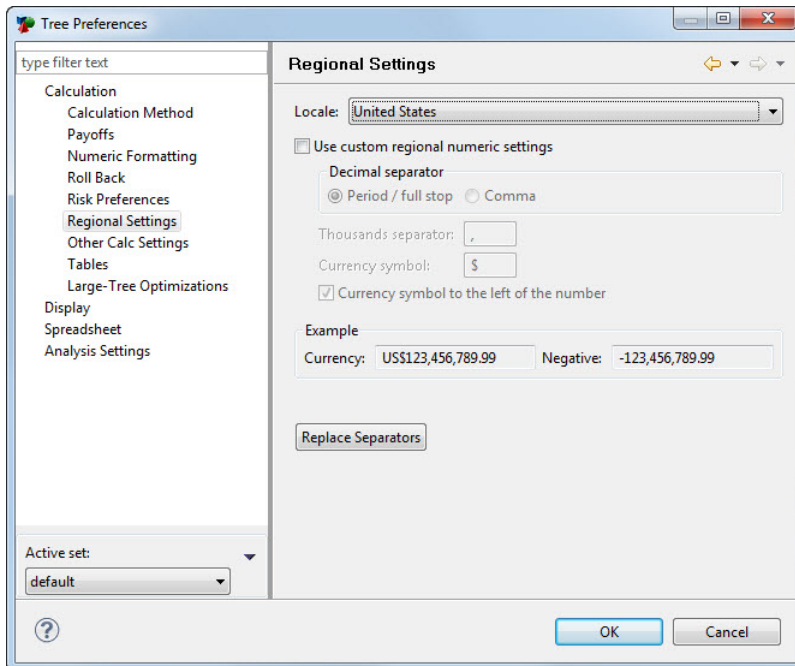
1375

The operating system’s regional settings are also, by default, used by TreeAge Pro to determine what decimal and thousand separator characters are used in displaying calculated values.

However, TreeAge Pro includes preferences that enable particular trees to override the operating system’s regional numeric settings, allowing you to specify which characters to use for decimal and thousands separators.

To enable custom regional numeric settings in a tree:

- Select the tree and press the F11 key to open the Tree Preferences dialog.
- Select the category Regional Settings.
- Check the box *Use custom regional numeric settings*.
- Modify the regional settings options as necessary.



Tree Preferences - Regional Settings

These settings will apply when this particular model is opened, modified, and analyzed on *any computer, no matter what the operating system's regional numeric settings*.

These settings are particularly useful when sharing models with colleagues in other countries.

10.5 Overriding the optimal path at decision nodes

TreeAge Pro selects an optimal path at decision nodes based on the calculation method preferences covered in previous chapters.

At a selected decision nodes, you can force TreeAge Pro to apply the *opposite optimal path* criterion to that specified in the tree's preferences.

To change the optimal path for a given decision node:

- Select the decision node.
- Choose Node > Change Optimal Path from the menu.

The decision node will reappear with an arrow inside it. The arrow will point upward if that decision node has been reset for maximization, or downward if the node has been reset for minimization. If you reverse the optimal path for an entire tree in which the optimal path of one or more nodes has been individually changed, the arrows in all of the individually changed nodes will reverse, signifying that they remain different from the rest of the tree.

To revert back to the original optimal path selection, simply repeat the steps above.

11. Selecting Subtrees and Multiple Nodes

While some tree-building tasks require that a single node be selected, other operations can, or must, be performed on multiple nodes. This chapter describes the different methods used to select multiple nodes.

11.1 Selecting a subtree

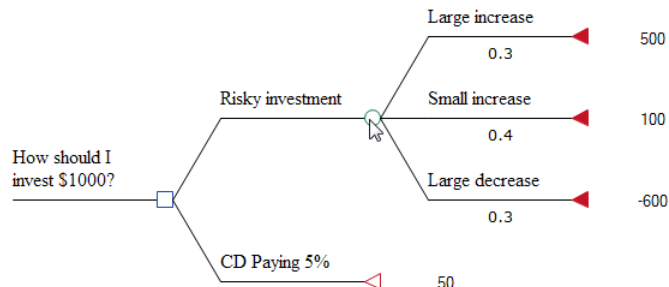
In decision trees, a *subtree* refers to a part of a tree. A node's subtree is comprised of all nodes and branches to its right (its “descendants”), but *not including the original node* (the subtree's root).

A few tree-building operations — copying, pasting, cutting, clearing, and cloning — require a special method for selecting a *subtree*.

To select a subtree:

- While holding down the *CONTROL* key, click on the subtree's root node.
- ... OR ...
- Right-click on the subtree's root node and choose "Select Subtree" from the context menu.

The figure below shows the result of selecting the *Risky Investment* node's subtree. Note that the *Risky Investment* node itself is not selected.



Select subtree



The methods of selecting multiple, individual nodes described in the next sections cannot be used to select a subtree.

11.2 Selecting multiple, unrelated nodes

TreeAge Pro offers several methods for selecting multiple, unrelated nodes. These techniques can be used when changing node types, entering payoffs, and a few other tasks.

To select a set of unrelated nodes:

- Select any node.

- While holding down the *SHIFT* key, select another node by clicking on it. Continue adding to the selection using the same shift-clicking operation.

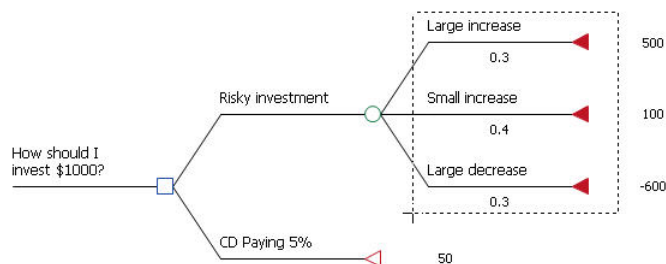
To remove nodes from the selection:

- Hold down the *SHIFT* key and click, one after the other, on each of the selected nodes that you wish to deselect.

It is possible to select multiple nodes by dragging a selection rectangle around them.

To select several adjacent nodes:

- Click and drag to create a selection rectangle that encloses the adjacent nodes you wish to select.
- Release the mouse button.



Select Multiple Adjacent Nodes



Note that you must enclose the entire node line and label for each node you want to select via the selection rectangle.

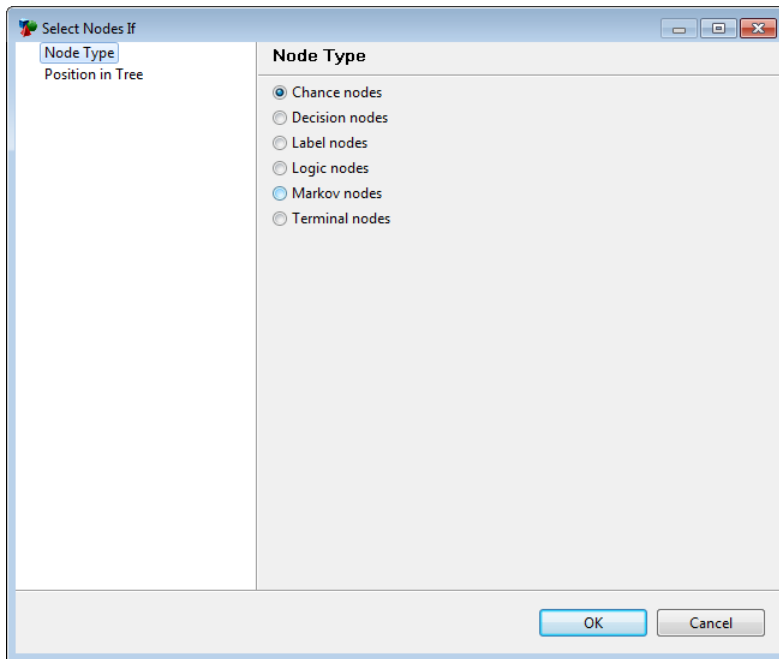
11.3 Selecting multiple nodes by characteristic

TreeAge Pro allows you to automatically select sets of nodes that share a node type or a position in the tree.

To select multiple nodes by characteristic:

- Choose Tree > Select Nodes If from the menu.
- Choose a selection method in the left pane of the Select Nodes If Dialog.
- Select the options associated with that method in the right pane of the Select Nodes If Dialog.
- Click OK.

This Select Nodes If Dialog is presented below.



Select Nodes If Dialog

12. Making Changes to Tree Structure

Building a tree is often a process of frequent revision. Also, new tree projects can often make use of subtrees from existing models. This chapter starts with a review of the basic tree building commands from the Decision Tree Tutorial Chapter, and then covers a set of features designed to help you move, remove, and duplicate parts of a tree (called “subtrees”).

12.1 A note on tree terminology

In tree models, familial terms are often used to help identify particular nodes and branches based on their relative locations.

- Nodes in its path back to the root node are its *ancestors*.
- Nodes in the paths to its right are its *descendants*.
- Branches of a *parent* node are its *children*.
- Branches of a parent node are also each other's *siblings*.

12.2 Tree-building commands – a review

Here is a quick review of some of the basic tree-building commands introduced in the Decision Tree Tutorial Chapter.

To add branches:

- Select a node.
- Drag a node from the Tree Diagram Editor Palette onto the Tree Diagram Editor. As you drag the new node onto the existing tree, it will appear in red in spots where the new node can be placed. Release the mouse when the new node is in the appropriate position in the model.
- ... OR ...
- Select a node and right-click on the decision node and select Add Branch from the context menu.

To delete a branch:

- Select the node.
- Choose Edit > Cut from the menu to cut the node from the model.
- ... OR ...
- Right-click on the node and choose Cut from the context menu.



Note that branches can also be cut by performing the control + X keyboard shortcut.

To change a node's type:

- Right-click on the node.

- Select Change Type > Terminal from the context menu (see figure below).
- ... OR ...
- Select a node with no branches.
- Click the "change node type" icon on the toolbar.
- Select the appropriate value from the list of node types.

12.3 Editing a single node vs. subtrees

Many revisions to tree structure can be accomplished by inserting, deleting, or moving one node/branch at a time, or by changing the order of branches emanating from a node.

The Cut, Copy, and Paste commands described later in this chapter can be used to efficiently carry out similar, but more complex, tree modifications – including inserting, deleting, and reordering *subtrees*, as well as duplicating nodes or subtrees.

12.4 Inserting nodes/branches

Nodes/branches can be inserted into a model in one of two ways - via the node context menu and via the Modeling Palette.



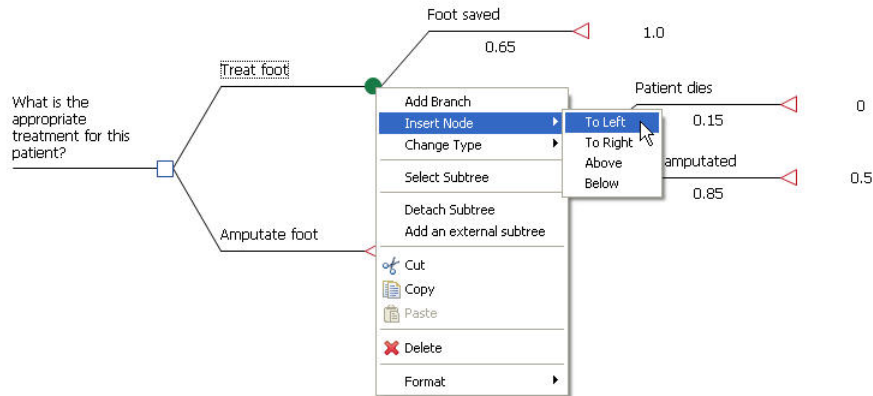
Note that inserted branches can be removed using the Undo command by selecting Edit > Undo from the menu or clicking *CONTROL + Z* on the keyboard.

12.4.1 Insert node via context menu

The Insert Branch options, in comparison to Add Branch, provide greater control over how and where new branches and nodes are created. In addition to the Add Branch functionality, Insert Branch can be used to add a sibling above or below a selected branch, or to make “generational” changes — i.e., inserting a node between a “parent” and its “children.”

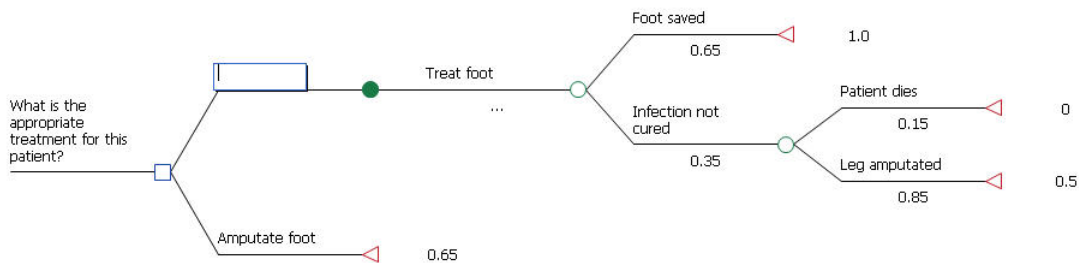
To insert a new node into a model via the context menu:

- Open the Get Started tutorial example tree, Rock Climber.
- Right-click on the the chance node "Treat foot".
- Select Insert Node > To Left from the context menu.



Insert branch

A new branch, ending in a chance node, is inserted between the root, decision node and the "Treat foot" node. See below.

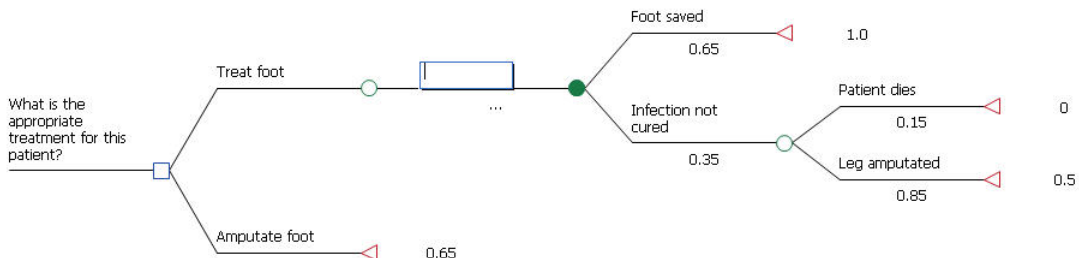


After inserting branch to left

Repeat the menu command, but try different directions to see what happens.

- Right-click on the the chance node "Treat foot".
- Select Insert Node > To Right from the context menu.

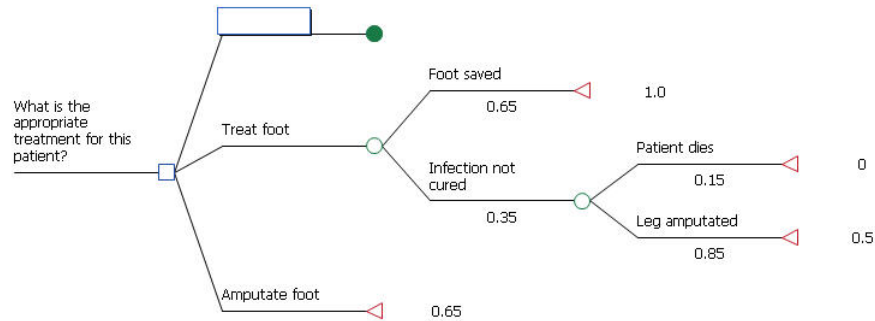
This time a new branch is inserted after the "Treat foot" node, whose existing branches are now attached to the new chance node. See below.



After inserting branch to right

- Right-click on the the chance node "Treat foot".
- Select Insert Node > Above from the context menu.

In this case, the new node is added as a higher sibling of the "Treat foot" branch. See below.



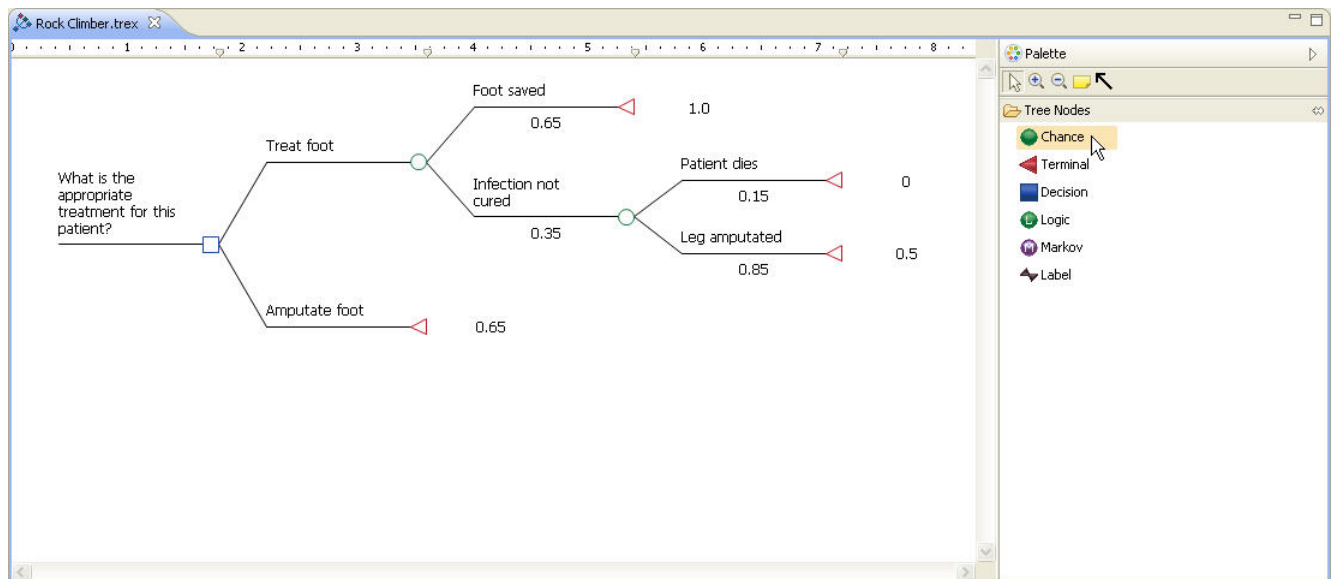
After inserting branch above

If a new node were inserted below the "Treat foot" node, the new node would be added as a lower sibling of the "Treat foot" branch.

12.4.2 Insert node via Modeling Palette

Nodes can also be inserted into a model by dragging a new node from the Modelling Palette into the Tree Diagram Editor.

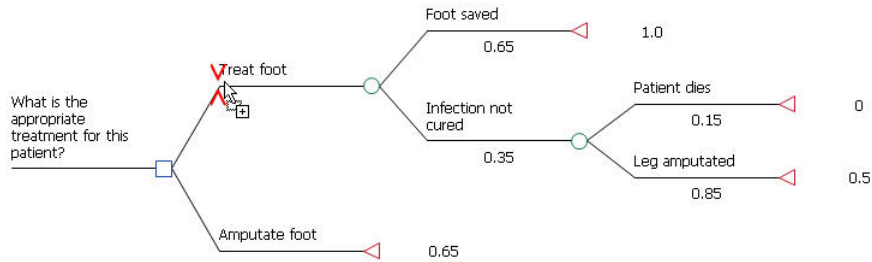
First select the node type you wish to insert from within the options in the Modelling Palette. See below.



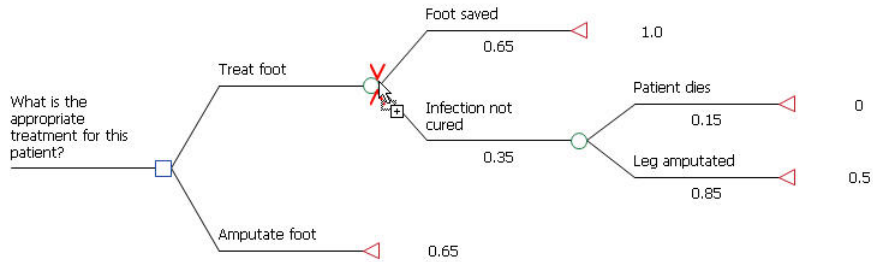
Select node type in Modelling Palette

Then drag the new node into the model where you want the new node inserted. Release the mouse button when the node is in the correct place.

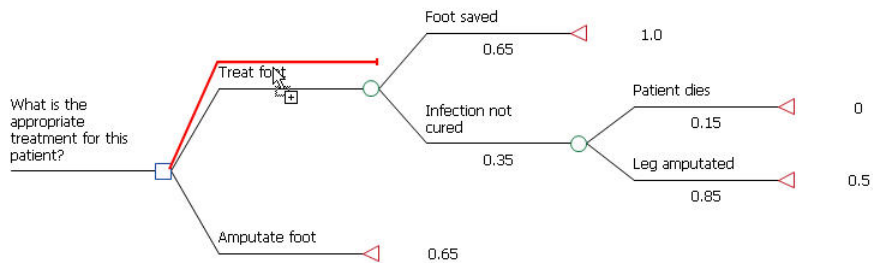
The four figures below show how to insert a new node to the left, right, above, and below the "Treat foot" node.



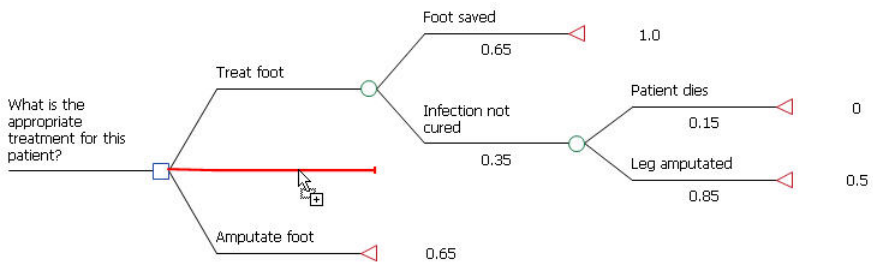
Insert/drag node to the left



Insert/drag node to the right



Insert/drag node above



Insert/drag node below

12.5 Moving/reordering nodes

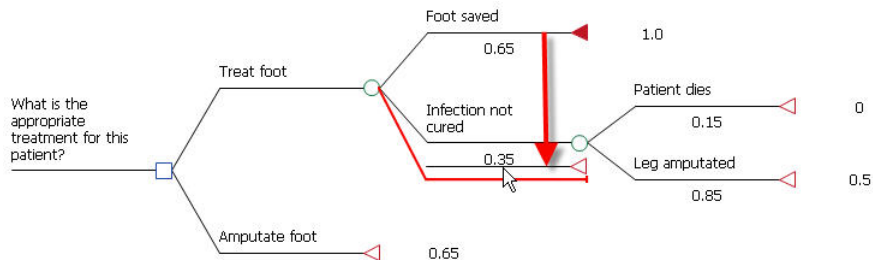
In prior versions of TreeAge Pro, a separate dialog was used to reorder branches. However, starting with TreeAge Pro 2011, moving/reordering nodes is handled by drag and drop functionality within the Tree Diagram Editor.

The prior section described how to insert nodes by dragging new nodes from the Modelling Palette. The same drag and drop options can be used with nodes that already exist within the model.

To move a node within a model:

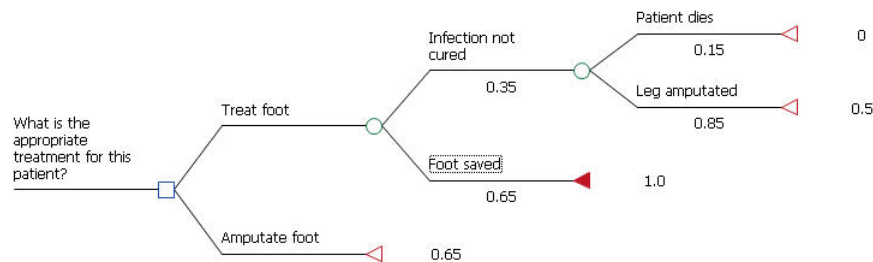
- Click on an existing node.
- Drag it to a new location within the model.
- Release the mouse button when you reach the proper destination location.

For example, you can reorder the branches of the "Treat foot" node in the Rock Climber tree by dragging the "Foot saved" node below the "Infection not cured" node. See below.



Move/reorder node

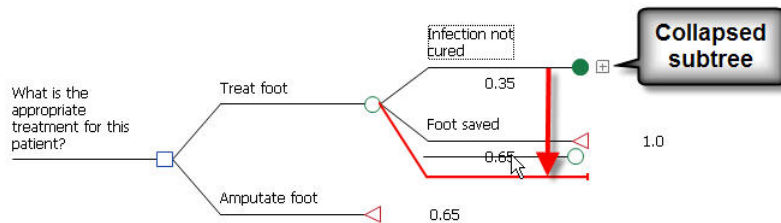
After dragging the node, the branches are reordered. See below.



After move/reorder node

Be careful moving nodes that are not endnodes (furthest to the right). For example, if you move the "Infection not cured" node back below the "Foot saved" node, the subtree anchored at the "Infection not cured" node will not automatically move with the node.

To move a node and its subtree together, first collapse the subtree, then move the node (see below). Alternatively, you can select the node and *all* the nodes in the subtree, then move the node to its new location.



Move/reorder node with subtree

12.6 Deleting nodes/branches

You can delete any node from the model.

To delete a node:

- Select the node.
- Right-click on the node and select Delete from the context menu.
- ... OR ...
- Choose Node > Delete from the menu.



Refer to the next section for information on deleting/cutting nodes that are not end nodes.

12.7 Cut, copy, and paste nodes and subtrees

Frequently, sections of a tree can be reused in another part of the same tree, or a different tree that you are working on. TreeAge Pro allows you to select a node, a set of nodes, or an entire subtree, copy or cut it, and then paste it to one or more nodes in any open tree.

In addition to being able to duplicate and move subtrees in the Tree Diagram Editor, it is also possible to manipulate subtrees using the mouse and **CONTROL** key in the Tree Explorer View. Refer to the Tools and Functions for Complex Trees Chapter for details on working in the tree explorer.



Besides text, tree nodes, and subtrees, other items that can be copied and pasted in TreeAge Pro include dependency diagram nodes, variable definitions, and links with Excel. Items that can be copied from, but not pasted into, TreeAge Pro include text reports, model images, and graph images.

12.7.1 Cut/copy/paste a single node

The Cut, Copy, and Paste commands are available for any node or set of nodes.

To copy a node:

- Select a node.
- Select Edit > Copy from the menu.

- ... OR ...
- Click *CONTROL* + C on the keyboard.
- ... OR ...
- Right-click on the node and select Copy from the context menu.

Once copied to the clipboard, the node can be pasted to any node as a new branch. It can be pasted to any non-terminal node (branches not allowed) in the same tree or a different one. All aspects of the node (i.e., probability, payoff, variable definitions, etc.) are included with the copy.

To paste a copied node:

- Select any single node that is not a terminal node.
- Select Edit > Paste from the menu.
- ... OR ...
- Click *CONTROL* + V on the keyboard.
- ... OR ...
- Right-click on the destination node and select Paste from the context menu.

After being pasted, the copied node remains in the clipboard, and can be pasted at additional locations. It will remain in the clipboard until something else replaces it in the clipboard — including from an application other than TreeAge Pro.

It is also possible to cut a node — i.e., remove it from its current location, in order to paste it in a new location.

To cut a node:

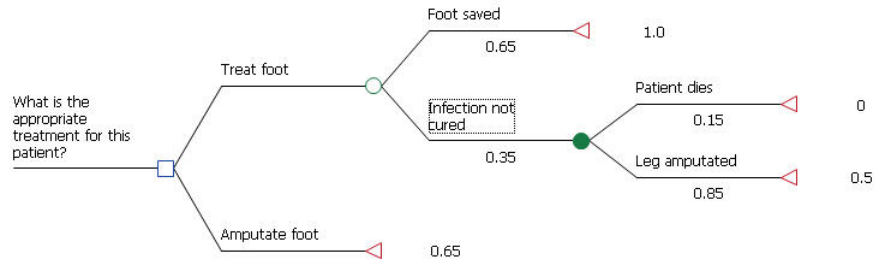
- Select a node.
- Select Edit > Cut from the menu.
- ... OR ...
- Click *CONTROL* + X on the keyboard.
- ... OR ...
- Right-click on the node and select Cut from the context menu.



Special care must be taken when cutting and/or pasting anything other than a single endnode (i.e., any node without branches, except the root node). These actions can sometimes cause unexpected shifts in the model structure.

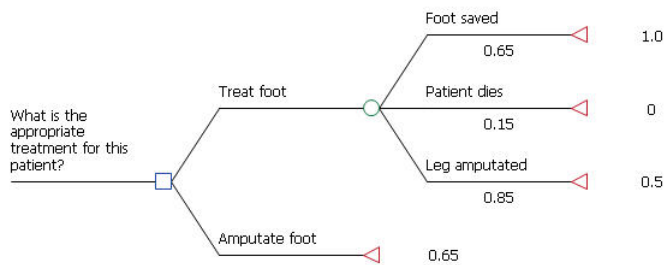
Note that you can always use the Undo command (*CONTROL* + Z) to undo the last changes to the model.

Consider what happens if you cut a node that is not an endnode.



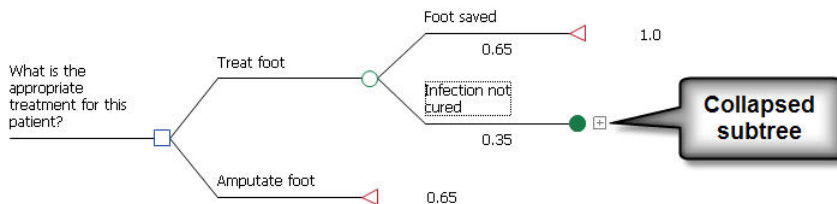
Cut non-endnode

If you cut the node "Infection not cured" in the model above, the cut node's branches all shift to the left to become descendents of the original node's parent, in this case the "Treat foot" node. See below.

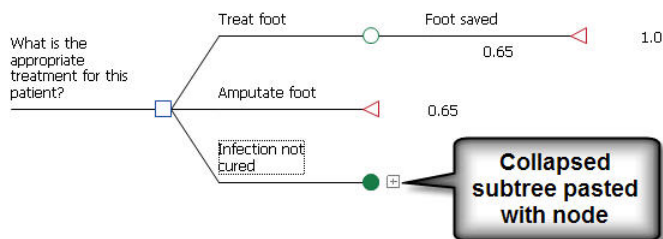


After cutting non-endnode

The chance node "Infection not cured" can then be pasted elsewhere, but its descendents are not pasted with it. However, if you first collapse the subtree (Node > Collapse Subtree in the menu), then the subtree is copied with the node.



Cut node with collapsed subtree



Collapsed subtree pasted with node

12.7.2 Cut/copy/paste a subtree

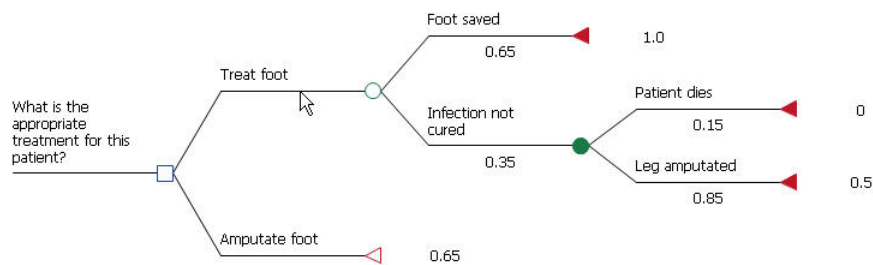
Duplicating large tree structures, rather than building each one from scratch, can both save time and avoid errors. In many trees, some alternatives will be structural identical or closely similar, differing perhaps in particular probabilities or payoffs.

Before a subtree can be copied, it must be selected.

To select a subtree:

- Right-click on a single node that has branches attached (i.e., not an endnode) and choose "Select Subtree" from the context menu.
- ... OR ...
- Hold down the *CONTROL* key and click on a single node that has branches.

When the subtree is selected, all the descendents of the "subtree root" will be selected while the "subtree root" itself will not be selected as it is not considered part of the subtree itself. The collection of nodes in the selected subtree can then be copied or cut. In the figure below, the subtree emanating from the "Treat foot" node was selected.



Select subtree

Once a node's subtree is selected, the Copy Subtree command becomes available.

To copy a subtree:

- Choose Edit > Copy from the menu.
- ... OR ...
- Click *CONTROL* + C on the keyboard.

Once copied to the clipboard, the subtree can be pasted to any non-terminal node — one with or without branches, in the same tree or a different one.

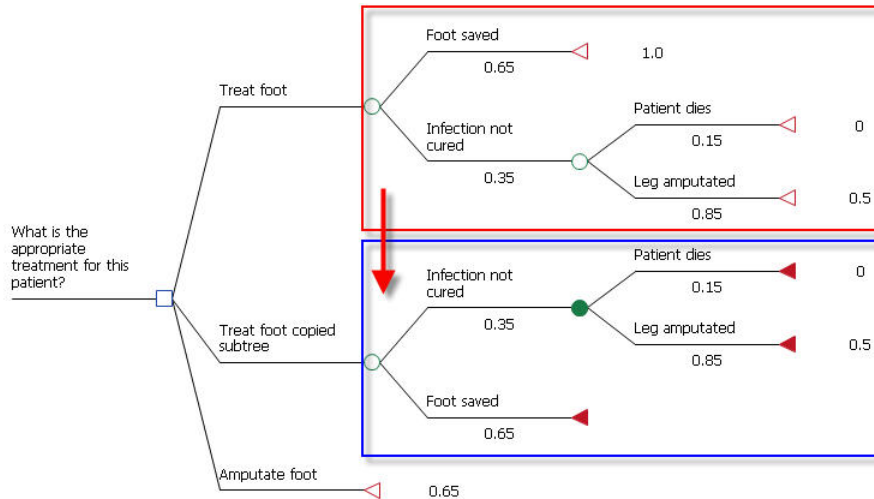
To paste a subtree:

- Right-click on any single non-terminal node and choose Paste from the context menu.
- ... OR ...
- Select any single non-terminal node and click *CONTROL* + V on the keyboard.



If you want to carry all information (e.g., Markov, DES) from the source subtree to the pasted subtree, be sure the source and destination subtree root nodes have the same node type.

In the figure below, the "Treat foot" subtree was copied to the new node "Treat foot copied subtree".



After pasting subtree



Note that the node "Treat foot copied subtree" above was not created by the copy and paste subtree functions. That node was created separately.

To move a subtree, rather than duplicate it, use the Cut Subtree commands.

To cut a subtree:

- Select a subtree as specified above.
- Choose Edit > Cut from the menu.
- ... OR ...
- Click *CONTROL* + X on the keyboard.

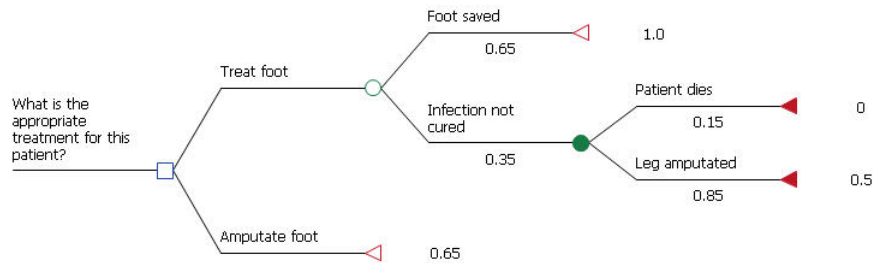
This will cause the subtree (but not the subtree's root) to be removed from the tree and placed on the clipboard. As with a copied subtree, when a subtree is cut to the clipboard, the Paste Subtree command becomes available.

12.7.3 Cut/copy/paste multiple nodes

When a subtree is cut/copied/pasted, the structure of the subtree is maintained to preserve the parent/child relationships among the nodes. When an assortment of nodes are selected, there may or may not be parent/child relationships among the nodes. Therefore, special care must be taken to ensure that the cut/copy/paste functions change the model as desired.

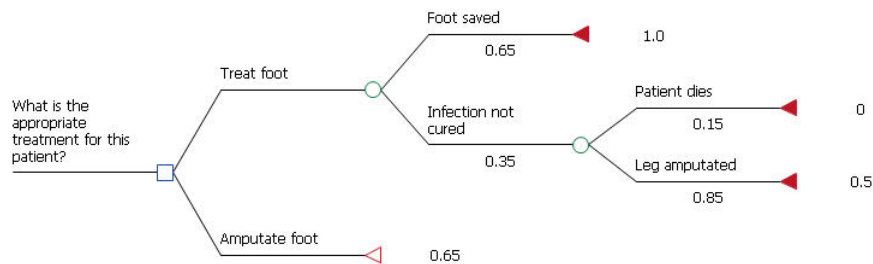
For example, if you select the three nodes as shown below, there is a parent/child relationship among the nodes. Therefore, those nodes could be copied and pasted to another location while maintaining

those relationships in the pasted destination (i.e., the nodes "Patient dies" and "Leg amputated" will remain children of the node "Infection not cured").



Select multiple nodes with parent/child relationship

However, in the next figure, there is no parent-child relationship among the selected nodes. Therefore, the selected nodes would all be pasted as branches of the destination node.



Select multiple nodes without parent/child relationship

It is also possible to cut/copy/paste a set of nodes where there are parent/child relationships among some of the nodes and not among others. In such a case, existing parent/child relationships will still be maintained.

12.8 Cut, copy, and paste text

It is possible to cut, copy, paste, and delete text in node descriptions, as well as in probabilities, payoffs, and any other formulas or values you type in TreeAge Pro.

Before cutting, copying, or deleting text, you must select the targeted letters, numbers, or words.

To select individual characters of text or formulas:

- Either use the mouse to: a) click and drag from one end of the desired selection to the other; or b) click in front of one end of the selection, and then shift-click after the other end.

To select one or more words of text (or a formula):

- Use the mouse to either: a) double-click on the first word of the selection and drag to the last word; or b) double-click on the first word of the selection, and then shift-double-click on the last word.

Once the text is selected, then you can choose the appropriate command from the Edit menu.

To copy (duplicate) selected text:

- Choose Edit > Copy.
- ... OR ...
- Click *CONTROL* + C on the keyboard.

To cut (move) selected text:

- Choose Edit > Cut.
- ... OR ...
- Click *CONTROL* + X on the keyboard.

The Copy and Cut commands both place the selected text on the clipboard. The text on the clipboard can then be pasted into any TreeAge document, or into another program.



The Cut, Copy, and Paste commands can also be accessed by right-clicking on the selected text and choosing the command from the context menu.

The Paste Text command can be used to insert text or formulas both place the selected text on the clipboard. This text can then either be pasted into any TreeAge document, or into another program.

To paste (insert) text:

- Place the text cursor in the desired location.
- Choose Edit > Paste.
- ... OR ...
- Click *CONTROL* + V on the keyboard.

Selected text can be deleted without placing it on the clipboard by pressing the *DELETE* key or the *BACKSPACE* key.

12.9 Multiple clipboards

TreeAge Pro has four tree clipboards, where one is active at any one time. This means, in effect, that you can cut or copy subtree X without losing subtree Y that is currently on the active tree clipboard.

Prior to cutting or copying another subtree, simply activate one of the empty tree clipboard's by selecting it in the Edit menu.

To select a different clipboard:

- Choose Edit > Tree Clipboard > Clipboard # from the menu.

12.10 Undo and Redo

Each tree (and dependency diagram) retains in memory details of the last actions that you took. Beginning with the most recent action and working back one action at a time, you can remove most all changes to structure, values, preferences, etc.



Note that the Undo “chain” is broken by some analysis and tree-building commands – TreeAge Pro will usually warn you also if a command cannot be undone.

To undo a change to your model:

- Choose Edit > Undo.
- ... OR ...
- Click *CONTROL* + Z on the keyboard.

If you go back too far in undoing modifications to the model, you can use the Redo command, also found in the Edit menu. The Redo command essentially allows you to undo previous Undo action(s).

To redo an undone change to your model:

- Choose Edit > Undo [action].
- ... OR ...
- Click *CONTROL* + *SHIFT* + Z on the keyboard.

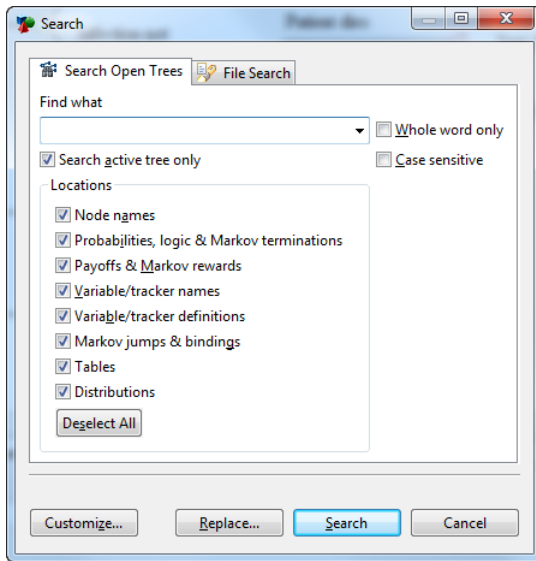
12.11 Find and replace text, formulas, and values

You can quickly search for and modify text and formulas at nodes in a tree using the Find/Replace tool. With some exceptions, this is often the best way to fix the spelling of a word used frequently in a tree, or to change a formula that appears in many payoffs, definitions, or other expressions. It is also often useful for simply finding particular nodes.

12.11.1 The Search Tree Dialog

To open the Search Tree dialog:

- Choose Tree > Find from the menu.
- ... OR ...
- Click *CONTROL* + F on the keyboard.
- The search dialog will default to "Search Open Trees" tab with "Search active tree only" checked. This will search only the currently active model.



Search Tree Dialog

The Search Tree dialog allows you to find and/or replace specific text in certain portions of the model or throughout the entire model.

12.11.2 Find Text

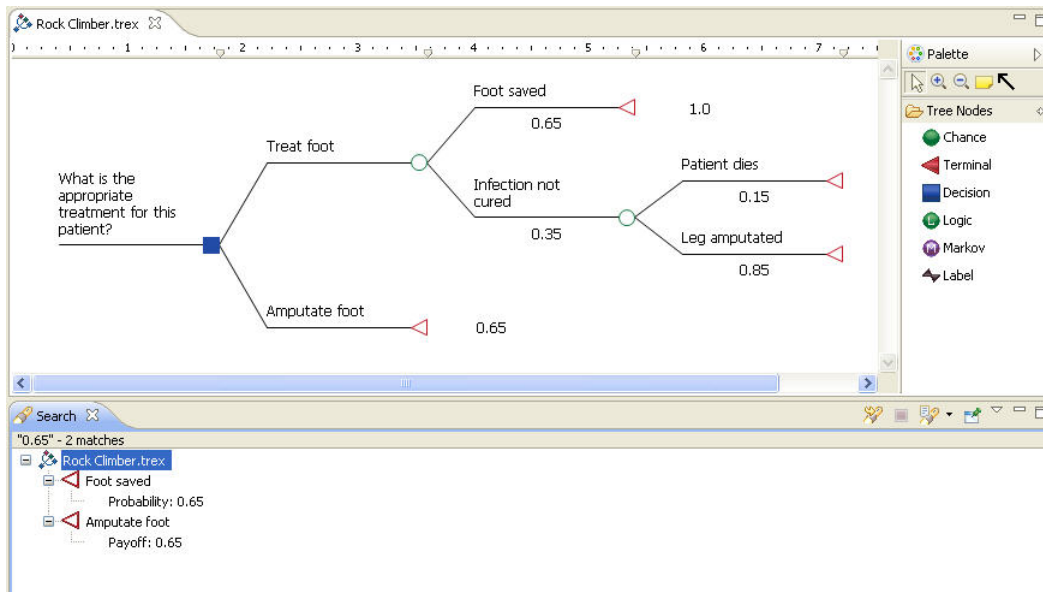
To find text in a tree:

- Open the Search Tree dialog.
- Enter text in the box labeled "Find what".
- Choose the matching options ("Whole word only" and "Case sensitive").
- Choose one or more search location(s).
- Click the Search button.

If you check the "Whole word only" option, TreeAge Pro will not search for partial word matches. If "Case sensitive" is checked, matching text must have the same combination of upper and lower case letters as the search text that you specify.

The locations refer to portions of the tree which should be searched for the search text.

The search results are then presented in the Search View.



Search Tree View

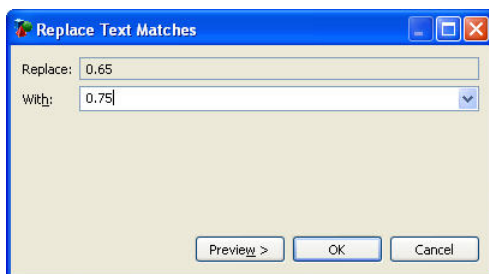
In the above example, the search text was "0.65". The nodes that include that text are presented in the Search View. Each node can be expanded/collapsed. When expanded, the node property or properties that contain the search text are displayed. If you double-click on the node (or its property), the node will be selected in the Tree Diagram Editor.

12.11.3 Replace Text

To replace text in a tree:

- Open the Search Tree dialog.
- Enter text in the box labeled "Find what".
- Choose the matching options ("Whole word only" and "Case sensitive").
- Select one or more search location(s).
- Click the Replace... button.

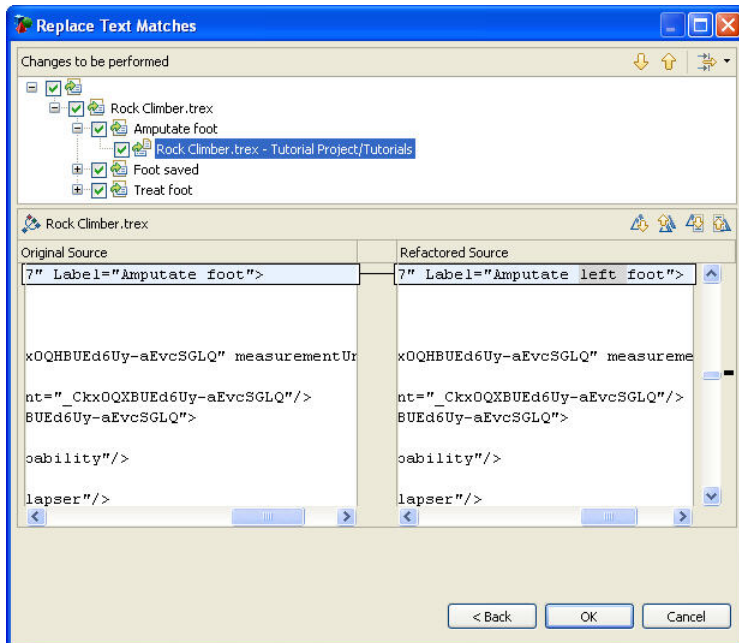
The Replace Text Matches dialog will open to prompt you to enter the replace text.



Replace Text Matches Dialog

You then have two options for replacing text. If you click OK, all occurrences of the Search Text that fit both the matching options and search locations will be replaced.

If you click Preview, then a Replace Text Matches preview dialog shows all replacements that could be made by this action.



Replace Text Matches Preview Dialog

The Replace Text Matches preview dialog allows you to examine all the changes that would be made to the model, including the differences in the file structures between the "before change" and "after change" file states. You can select which of the changes you want made, then click OK to process the changes.



The Replace command is not recommended for changing a variable's name. Instead, edit the variable's properties in the Variables and Tables list.

12.12 Using the Probability Wheel View

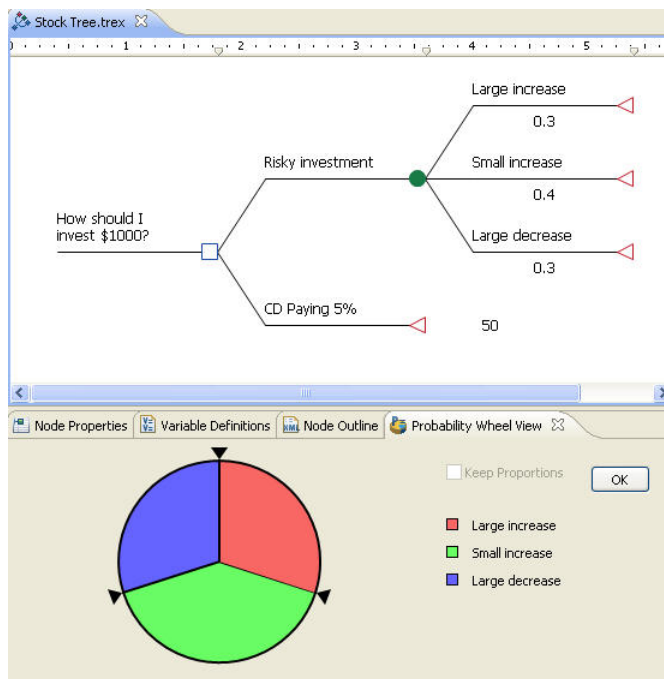
A frequent problem encountered in decision analysis is the assignment of subjective probability assessments to chance events. Many people find it easier to use a graphical aid in assigning probabilities. One tool designed for this task is a probability wheel. TreeAge Pro provides this functionality through the Probability Wheel View.

The Probability Wheel View is a tool for assigning probabilities to each branch of a selected Chance node. The Probability Wheel View works whether or not you have already assigned probabilities to the branches. If you have assigned probabilities, they will be used as initial values for the wheel. Each branch will be assigned its own colored portion of the wheel.

As an example, the probability wheel could be used to aid in assigning relative weights to the two outcomes of the risky investment in the Get Started tutorial example tree, Stock Tree, used in earlier chapters.

To open the Probability Wheel View:

- Select a chance node with branches.
- Choose Views > Probability Wheel from the toolbar.




Probability Wheel View with Tree

The probability wheel shows sections of the wheel to represent the probability associated with each branch. Drag the pointers around the edge of the wheel until the sizes of the section match your best assessment of the relative likelihood of outcomes.

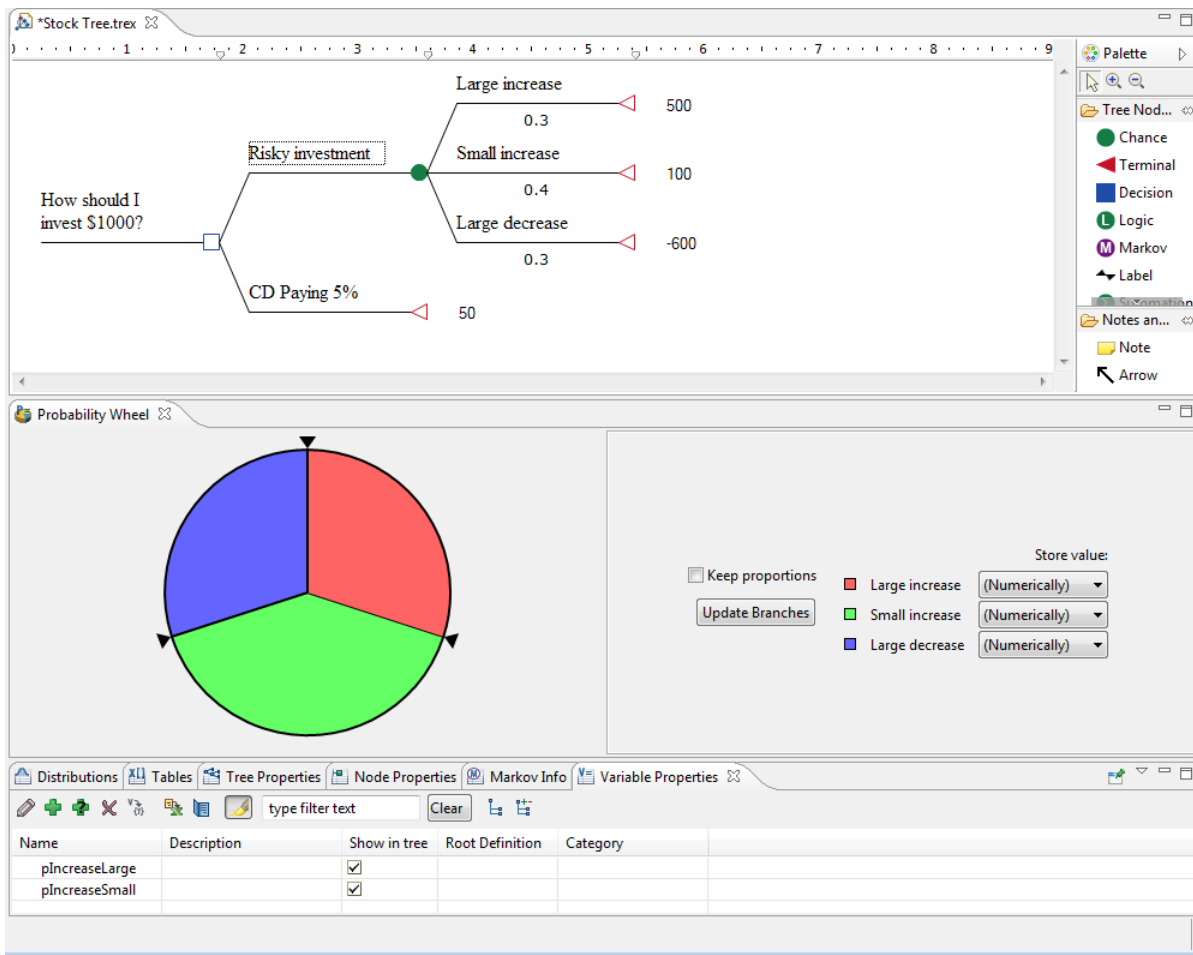
If the selected node has three or more branches, you will see a check box named "Keep proportions". If checked, the probability bounded on the right side by the selected marker is changed independently, while ratios are maintained for the remaining probabilities.

Click the mouse button on any wedge to display the numeric value (probability) of that wedge. Right-clicking will display that wedge's starting value.

Click the OK button to apply the probabilities from the Wheel to the branches of the selected Chance node. If you don't want to use the new branch probability values, just close the Probability Wheel View or select another element in the Tree Diagram Editor.

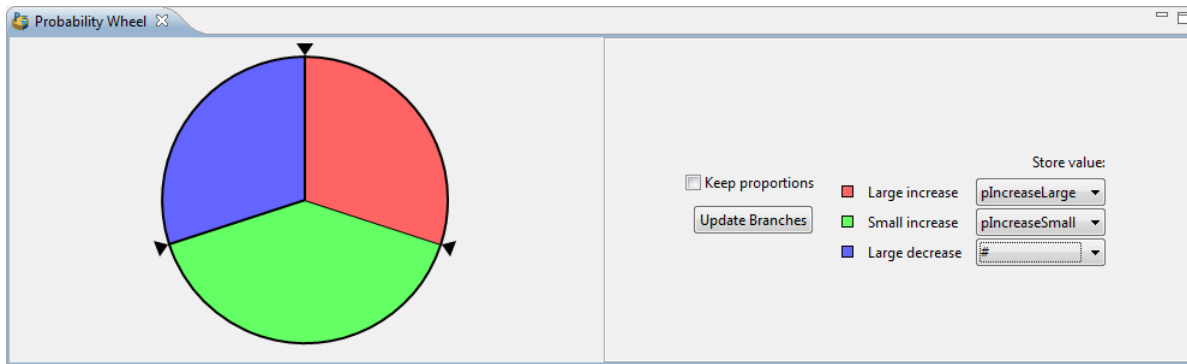
 The probability wheel should only be used to edit probabilities that are entered as numerical expressions. Other expressions can be converted to numerical expressions within the probability wheel to allow for editing.

Note that it is also possible to store the probability values from the probability wheel in variable definitions. To do so, you must first create the variable(s) via the Variable Properties View. After adding two variables *pIncreaseLarge* and *pIncreaseSmall* to the model and clicking off and back onto the *Risky Investment* node, additional options are shown in the Probability Wheel View. See the figure below with both the Probability Wheel and Variable Properties View displayed.



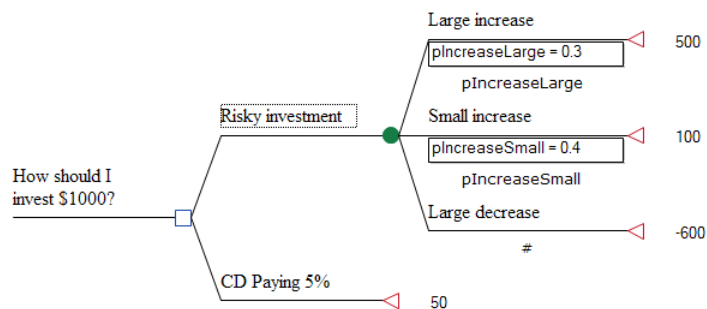
Probability Wheel before variables selected for storage

The new variables can now be selected for storage of the probability values from the wheel as shown below. Note that the complement "#" was selected for the third branch.



Probability Wheel with variables selected for storage

After clicking the Update Branches button, the variables are defined at the specific branches with the value from the probability wheel as shown below.



Variable definitions created by Probability Wheel

You might choose to move those variable definitions to the root node to make them accessible in other areas of a model.

13. Annotating the Tree

In addition to the basic ability to enter text labels at event nodes in the tree, TreeAge Pro includes a number of other options for annotating trees covered in this chapter.

The next chapter covers a variety of options for controlling the layout, formatting, and contents of decision trees.

13.1 Node label options

13.1.1 Text wrapping

Every node in a tree has a node label that is used to describe the purpose/function of a node. Entering node label text has been covered in earlier chapters. However, this section will describe how to control text wrapping of the node label.

In past versions of TreeAge Pro, node label text would not wrap unless you entered a carriage return (via the *ENTER* key). In TreeAge Pro 201x, the node label text wraps automatically based on the width of that node generation.

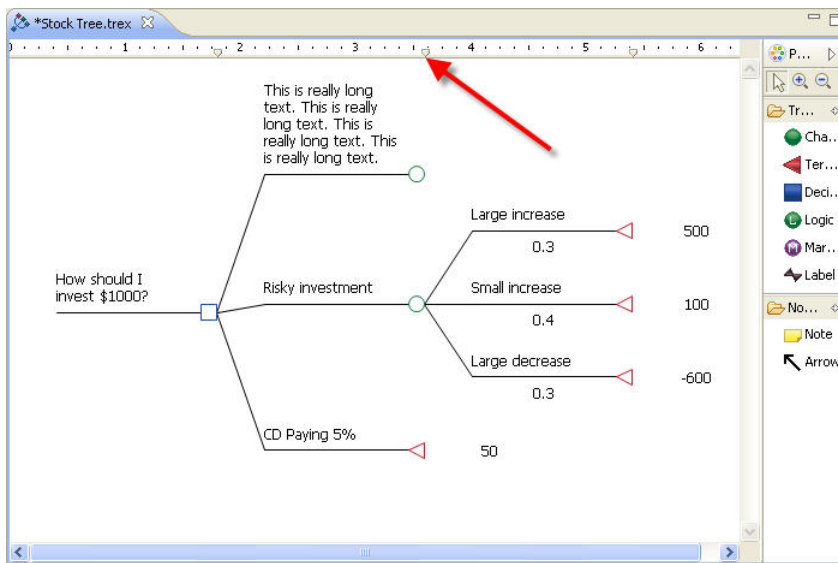
Also, in TreeAge Pro 201x, if you click the *ENTER* key when entering the node label, the currently entered node label text will be stored with the label and application control leaves the node label. You can still force wrapping at specific points in the text by clicking *CONTROL + ENTER* on the keyboard.



The keyboard command to force wrapping on a Mac is also *CONTROL + ENTER*.

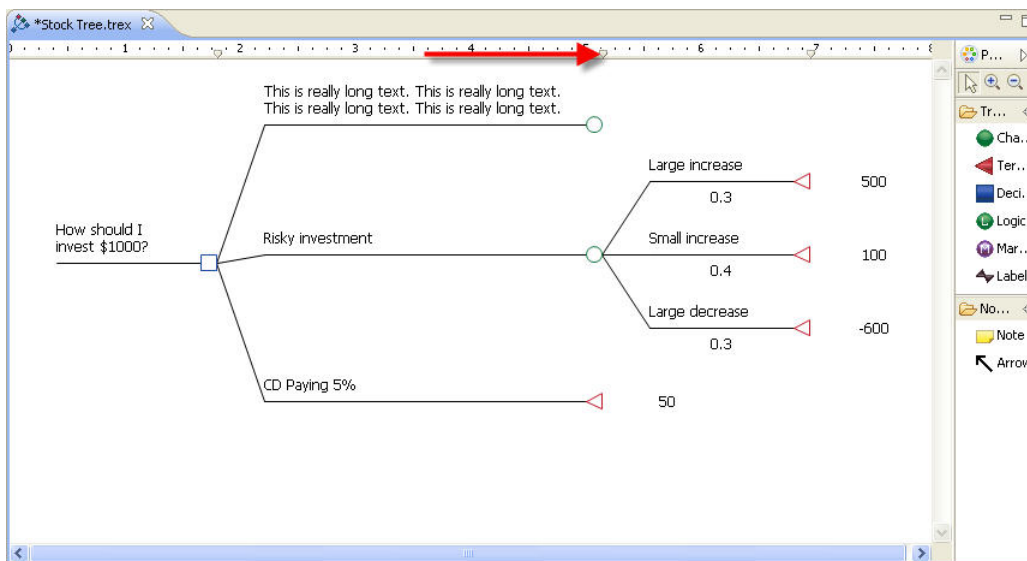
Since the node no longer stretches to accommodate the length of the node label text, you can now stretch a node generation via the ruler at the top of the Tree Diagram Editor.

In the figure below, the top node's label forces the text to wrap several times.



Node label text wrapped - before stretching node generation

Note the "tab pointer" in the ruler that is highlighted above. By dragging the "tab pointer" to the right, the entire node generation is stretched to make more room to display node label text, variable definitions, etc. See below.



Node label text wrapped - after stretching node generation



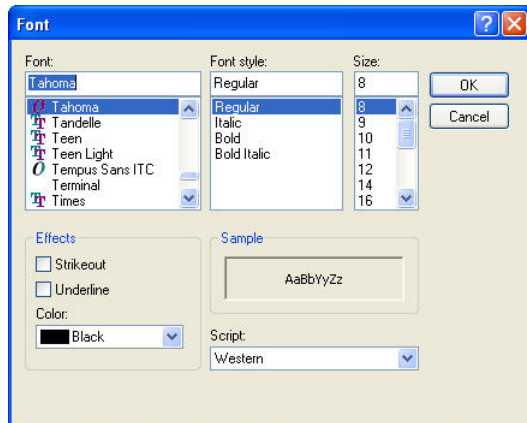
A node generation represents the set of nodes that are aligned vertically within the model. The ruler can be used to stretch a node generation, but it cannot be used to shrink the node generation beyond the standard minimum length.

13.1.2 Text formatting

TreeAge Pro supports multiple fonts, font styles and font sizes. The Fonts category of Tree preferences control the default text formatting for specific elements of a tree (e.g., node labels, probabilities, etc.). However, you can also set text formatting to a specific node label.

To change the text formatting for a specific node label:

- Right-click on the node and choose Format > Font from the context menu.
- Set the formatting options for the node label using the following Font dialog.



Font dialog

13.2 Label nodes

A label node, which uses a simple black “zigzag” as its symbol, acts like a placeholder. Label nodes have no impact on calculations, and cannot have more than one branch. A label node (or a series of them) can be inserted between event nodes to more clearly identify additional steps in a particular path.

To add a label node before an event node:

- Right-click on the event node and choose Insert Node > To Left from the context menu.
- Right-click on the new node and choose Change Type > Label from the context menu.

You could also change an existing endnode to a label node, and then use the Add Branch or Insert Node command to add the one allowed branch to the label node. A node with more than one branch cannot be changed to a label node.

For calculation purposes, a label node behaves like a decision node with one branch, or a chance node with one branch having a probability of 1.0. The value of the label node is simply the value of the node immediately to its right.



You can use a label node to separate dependent calculations. For example, you might have a tracker variable that is used in an expression, but the tracker needs to be assigned a new value before it is used in the expression. In such a case, you could insert a label node to the left of the node with the expression. Then you could assign the new value to the tracker in the label node. The tracker will then have the new calculated value when you reach the node with the expression the references that tracker.

13.3 Notes and arrows

Models often benefit from the use of longer explanations than are desirable in branch descriptions. Node comments, described in the previous section, are useful for the model builder, but not for someone looking at a printout or image of the model.

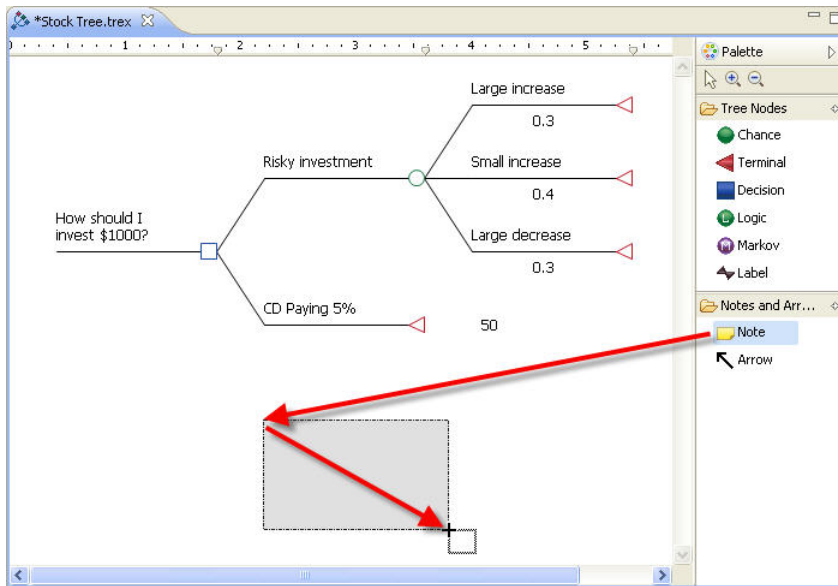
Using notes and arrows, you can provide the model's audience with an overview of the whole model, or specific nodes or subtrees.

13.3.1 Creating notes

Any number of notes can be created. Boxes are initially placed independent of any node, but can be bound to particular nodes if desired. Each box can use a different font, but all must use the same outline format (solid/dashed/none).

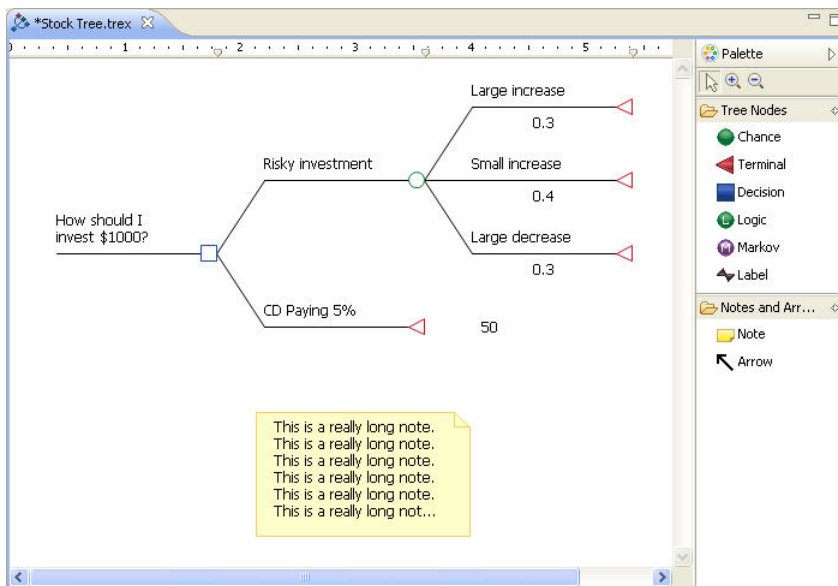
To draw a note in a tree:

- Click on Note in the Tree Diagram Editor Palette. The mouse cursor becomes crosshairs.
- With the mouse, click and drag somewhere in the Tree Diagram Editor. Make the box large enough to hold the text you wish to enter.



Create note

A yellow rectangular box is created that resembles a "Post-It" note. Within the box is a blinking text insertion caret, indicating that text can be entered in the note by typing. Text entered in the note will automatically wrap to the width of the box. If you enter text that does not fit into the box, the visible text will be truncated with an ellipsis (...) indicating that some text is not visible. Resize the note to display additional text (see next section).



After note is created and text is entered

The font of the active note can be changed. Each note can use a different font, but all text within a single note uses the same font.

To change the font of a note:

- First deselect the node by clicking elsewhere in the tree.
- Right-click near the edges of the note box (away from the text), and choose Format > Font from the context menu.
- Select the font options from the font dialog.

To see what the note will look like when printed, deselect it by clicking elsewhere in the tree.

The other options in the right-click context menu can be used to change the note's other visual characteristics, including text, fill, and line color.

To change the note outline:

- Right-click on the note and choose from the Format > Line Color or Line Type sub-menus.

13.3.2 Changing text in a note

To change the text in a note, you must first activate the text.

To activate the text in a note:

- Select the note box, and then press the F2 key.
- ... OR ...
- Click three times on the text in the note. (The first click selects the note shape, the second selects the text box within the note shape, and the third activates the text insertion caret within the text.)

Once the text is activated, you can modify the text by typing or using standard editing commands (BACKSPACE, DELETE, cut/copy/paste, etc.).

13.3.3 Moving, resizing, hiding and deleting notes

In order to move, resize, or delete a note, you must first select it. A note that is activated for text entry, displaying the blinking text caret, is *not* selected.

To select a note:

- Click once on the note.

The selected note will display a small square with "handles" at each of its four corners and on each of its sides. These handles are used to resize the note.

To resize a note:

- Move the mouse over one of the note's handles until the cursor appears as a double-sided resizing arrow.
- Click on the handle.

- Click and drag the handle to change the size and proportions of the note.

The corner handles allow you to resize both the height and width of the note, while the side handles only allow you to resize one or the other dimension.

To move a note:

- Click on the note and drag it to the desired location.

The right-click context menu can be used to cut, copy or delete a note.

To delete a note:

- Right-click on the note and select Cut or Delete from the context menu.

The context menu's Copy command can be used to copy a note into the clipboard. The main menu's Edit > Paste command can then be used to create a duplicate copy of the original note.

13.3.4 Displaying/Hiding all notes

When notes are placed in a model, it may sometimes be useful to hide all the notes which have been created.

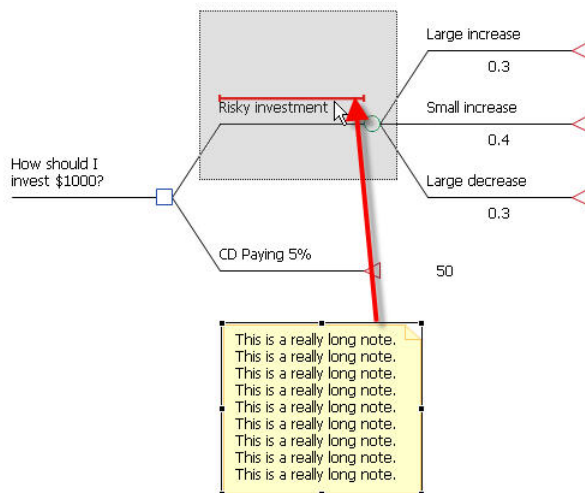
- Open Tree Preferences > Display > Node Text/Comments.
- Check the box "Hide all notes" to either hide/unhide all notes.

13.3.5 Binding a note to a node

When a note is placed in a model, it remains fixed in place relative to the top left corner of the tree window, and will not adapt to changes made to the tree. If nodes are added or deleted, for example, an existing note may overlap another object. This problem can be avoided by binding a note to a node.

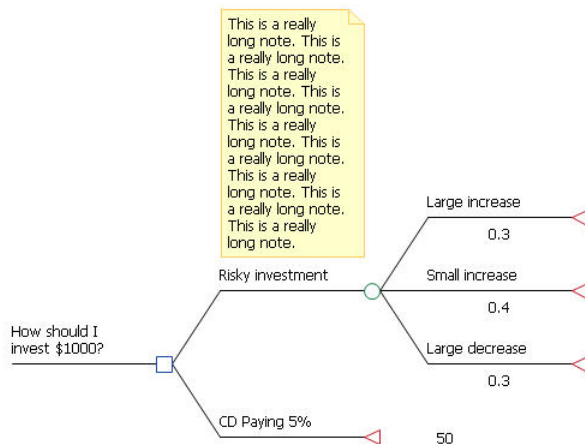
To bind a note to a node:

- Click on the note and drag it just above the node to which you want to bind your note.
- When the red line appears above the node, release the mouse button.



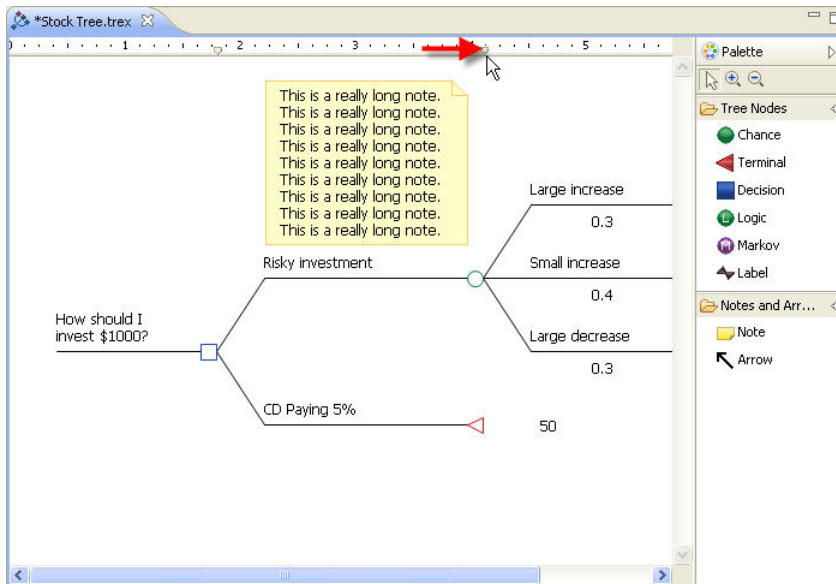
Bind note

The bound note will be aligned directly above the selected node, and will move with the node as the tree structure changes. The width of the note will be tied to the width of the bound node, and the text will automatically wrap to display all text.



After binding note

You can resize the node generation to better accommodate the contents of the bound note.



After resizing node generation for bound note

A bound note box can still be cut/deleted via the right-click context menu.

13.3.6 Creating arrows

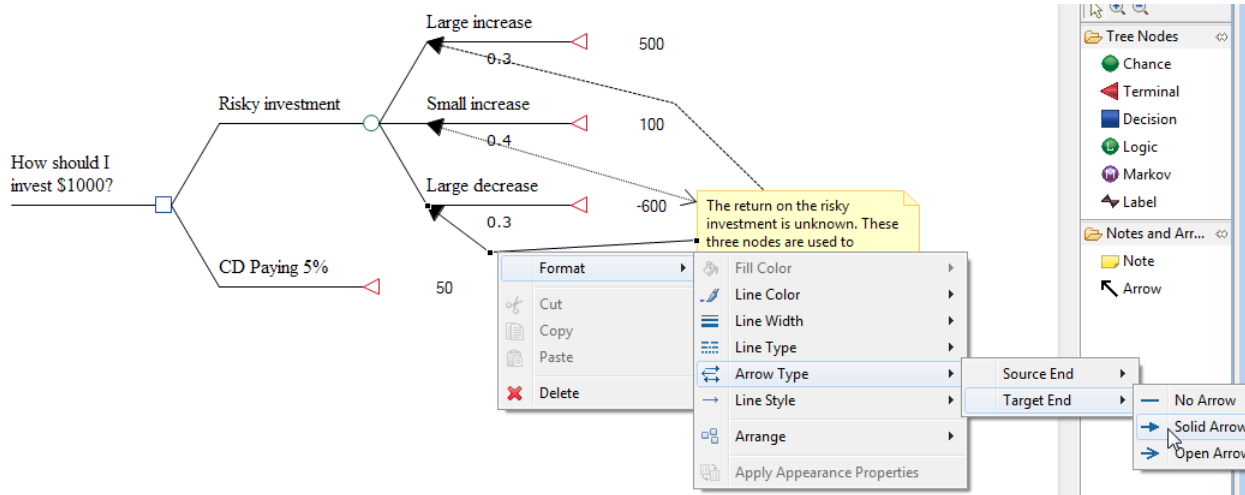
You can draw arrows in conjunction with annotating a tree. Arrows are generally used to associate an unbound note with one or more nodes.

To draw an arrow from a note to a node:

- Click on the Arrow in the Modeling Palette.
- Move the mouse to the note. When over the note, the cursor shape will be an arrow without the "not allowed" character.
- Click on the note and hold down the mouse button.
- Drag the mouse to the node.
- Release the mouse.

When the Arrow option is selected from the Modeling Palette, the mouse pointer will change to the shape of an arrow with a "not allowed" character. When the mouse is over an element in the Tree Diagram Editor that is valid for an arrow connector, the "not allowed" character will disappear.

By default, the arrow will be displayed as a dotted line with no arrow markers at either end. However, the arrow's format can be edited. The figure below shows a note with arrows to three nodes. The format of each arrow is different, and the formatting context menu (via right-click) is shown.



Arrows with formatting options

Arrows can be edited in the following ways.

- Use the formatting context menu to change the line color, width, type and style.
- Use the formatting context menu to change the arrow types for the source and target end.
- Use the formatting context menu to delete the arrow.
- Click on the arrow to create a bend point to break the arrow into separate line segments.
- Drag a bend point to form a straight line to remove it.

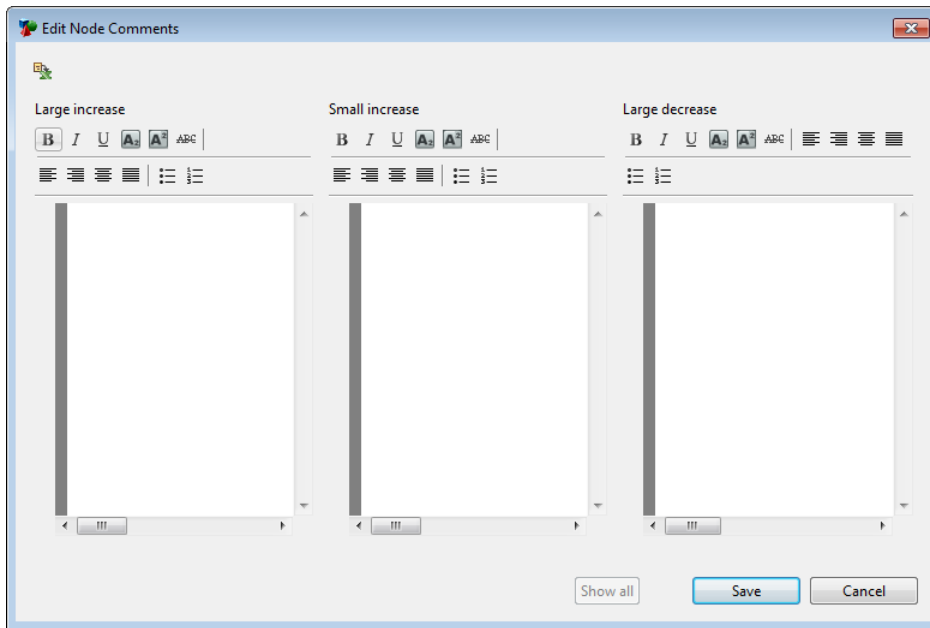
13.4 Node comments

Detailed comments can be assigned to the branches of a node and saved with the tree. Unlike notes and branch labels, the text of node comments is hidden.

Node comments are particularly useful for recording the basis on which probability assignments were made for the branches of a chance node.

To add node comments:

- Select a node that has at least one branch.
- Choose Node > Node Branch Comments from the menu.
- Enter node comments.
- Click Save.



Edit Node Comments Dialog

The dialog allows you to enter notes for each branch of the selected node. The notes can be formatted using the formatting toolbars. Formatted comments generate HTML output.

A red flag will appear above any node for which node comments have been entered. See below.

Node Properties View

Property	Value
Basic Info	
Label	Risky investment
Branch Comments	
Large increase	notes for large increase
Small increase	<html><body><P style="margin: 4;" align = "left">notes for small increase</P></body></html>
Large decrease	Notes for large decrease
Node Type	ChanceNode
Name ID	Node2
Clone Info	
Attach To Clone Master	0

Node comments shown in Tree Diagram Editor and Node Properties View

Node comments can also be seen in the Node Properties View. Click on the ellipsis button to edit the node comments for the selected branch. See above.

To remove node comments, delete the text in the Edit Node Comments Dialog.

14. Tree Display Preferences and Options

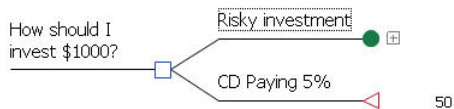
This chapter covers a wide variety of options for controlling the layout, formatting, and appearance of trees. Some of these features apply to calculated values displayed following roll back, while others affect display of the tree's structure.

14.1 Collapsing/hiding subtrees

The Collapse Subtree command can be used to temporarily hide any subtree (including the root node's subtree). It is particularly helpful when working with or presenting large trees.

To collapse a subtree:

- Select a node with visible branches.
- Choose Subtree > Collapse Subtree (or *CONTROL + J* on keyboard).



Collapsed subtree

The subtree emanating from the selected node is hidden and a plus sign (+) is displayed in its place, to the right of the node. The plus sign will also appear in place of the hidden subtree in printouts and exported images of the tree. Collapsing a subtree does not affect calculations.

14.1.1 Expanding hidden subtrees

Hidden subtrees can be uncollapsed in two ways: showing one generation of branches; or unhiding the entire subtree.

To expand a collapsed subtree one generation at a time:

- Select a node with a plus sign to the right.
- Choose Subtree > Expand Subtree Once (or *CONTROL + SHIFT + J* on keyboard).

The branches of the selected node will be displayed, but any subtrees attached to these branches will remain hidden and plus signs are displayed in their place.

To expand an entire collapsed subtree:

- Select a node with a plus sign to the right.
- Choose Subtree > Expand Subtree Once (or *CONTROL + ALT + J* on keyboard).



Visible clone copies (see Complex Trees Chapter) can be collapsed.

Collapsing a subtree in the tree explorer pane does not collapse the subtree in the regular tree view.

14.2 Aligning selected nodes

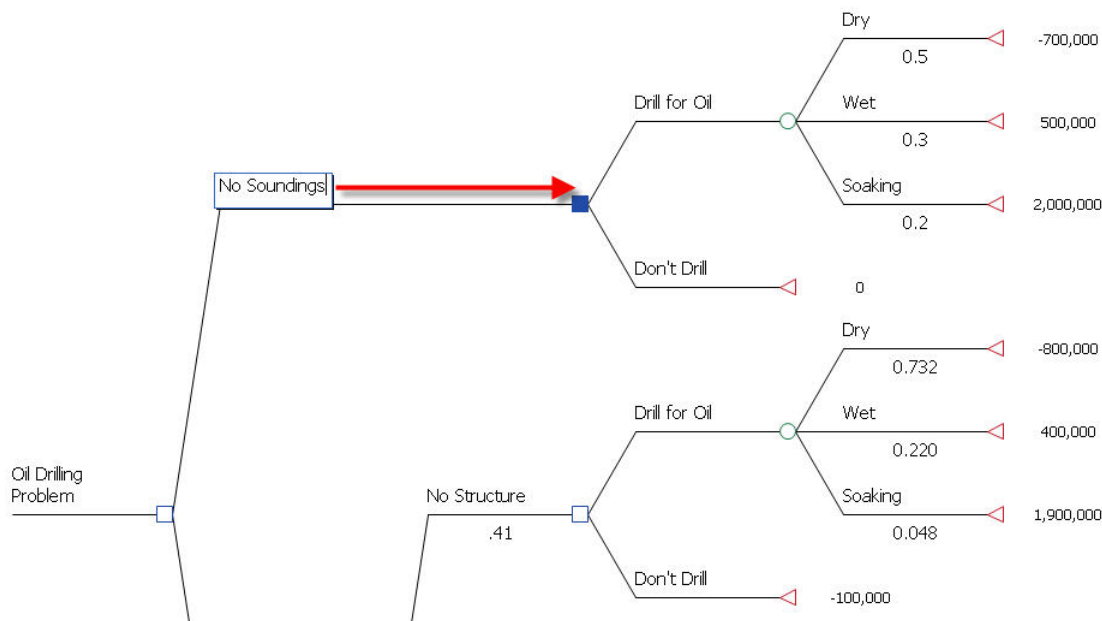
TreeAge Pro includes two options for adjusting the vertical alignment of nodes in a tree. This section deals with the use of the Skip Generation command to make manual adjustments to the vertical alignment of selected nodes. It is also possible to specify that all terminal nodes be vertically aligned automatically; this is covered in a later section, on tree display preferences.

In many trees, asymmetry in intervening events results in the related nodes not lining up vertically. It is sometimes desirable to force these related nodes to line up, resulting in a more intuitive layout of the tree.

To align a selected node with a node in a different subtree:

- Select a node.
- Choose Node > Skip Generation (or *CONTROL + J*).

This will extend the length of the selected node by one node generation and moves the selected node's subtree to the right.



Skip generation

Skipping one or more generations can be used to line nodes up vertically — for example, the multiple nodes representing a particular decision, in different paths. To skip more than one generation, simply select the menu command repeatedly.

To reverse the effects on the tree, simply un-skip generations at the node which skips a generation.

To remove an extra generation from the selected node:

- Select a node which is set to skip a generation.
- Choose Node > Unskip Generation (or *CONTROL* + *I*).

Skipping generations does not affect calculations.

14.3 Displaying terminal columns/roll back columns

When a tree is rolled back, TreeAge Pro normally displays next to each terminal node a roll back box containing the node's calculated payoff and, in an optimal path, its path probability. Previous chapters described some of the customizations that can be made to the roll back display.

Instead of showing the standard roll back boxes at end nodes, a tree can be set up to display user-defined columns of values to the right of visual end nodes during roll back.

Calculated values and other information that can be displayed in roll back columns include:

- payoffs, including extra (non-active) payoffs
- individual components of a complex payoff formula
- path probabilities
- scenario (i.e., terminal node) numbers

Roll back column options

Some important features of roll back columns include:

- During roll back, the table of values can be copied to a spreadsheet or other applications for reporting or further analysis.
- A row will be displayed for every *visual* end node, even those that are not terminal nodes — for example, if a node's subtree is collapsed (see above) or is a hidden clone copy (refer to the Complex Trees Chapter), a row is shown for the node at the base of the collapsed subtree.
- Calculated values can be displayed using custom numeric formatting.

Terminal column features

To add terminal columns in your model:

- Choose Tree > Tree Preference from the menu to open the Tree Preferences Dialog.
- Select the category Display > Terminal Columns.
- Check the "Show terminal columns" box.
- Click the "plus" button to add a terminal column.

Note that you can uncheck the "Show terminal columns" box later to return to regular payoff calculations without losing the terminal column preferences.

There are five columns in the terminal columns grid.

1. *Header*: Text to display above the column.
2. *Calculation*: Type of calculation to perform.
3. *Custom calculation*: Expression to calculate if Calculation value is "Custom".
4. *Numeric format*: Existing format option to apply to column.
5. *Custom*: Check to enter a custom format for this column

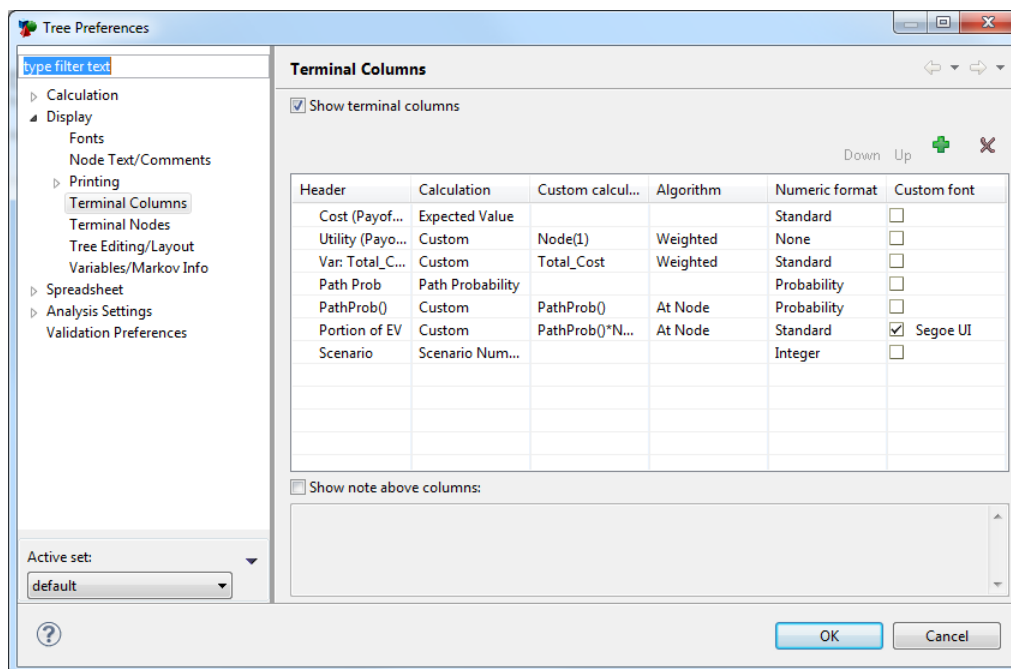
Columns in terminal columns grid

There are five Calculation options.

1. *Custom*: Enter any expression in the Custom calculation column.
2. *Expected Value*: The payoff value for the active payoff set.
3. *Incremental Value*: The difference in value between strategies.
4. *Path Probability*: The cumulative probability of reaching that terminal node. Only shows the recommended strategy.
5. *Scenario Number*: Counts from the top terminal node to the bottom.

Calculation options

The Special Features tutorial example tree "Terminal Columns" illustrates the use of terminal columns. Note the terminal columns from the tree preferences.



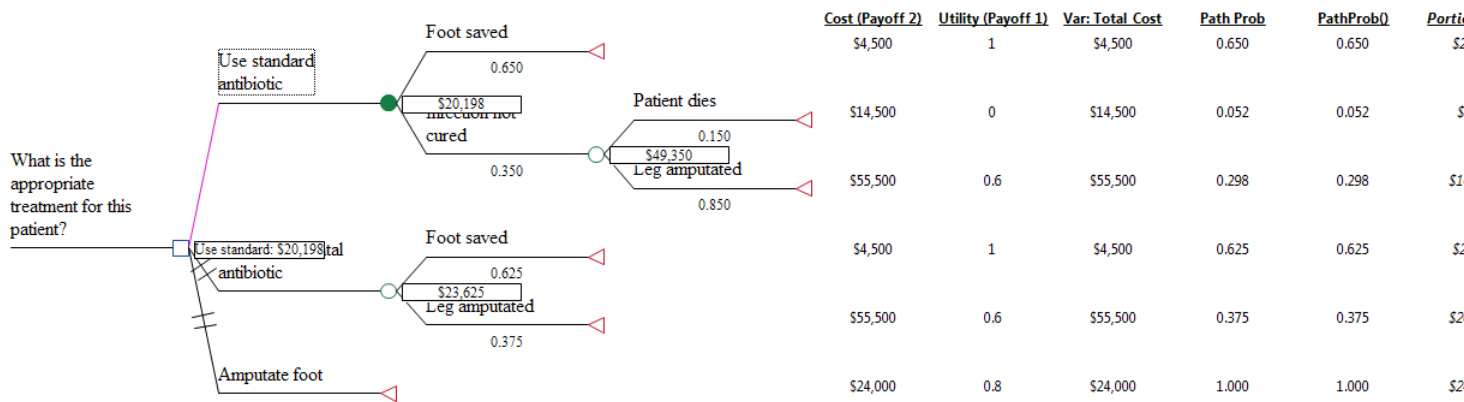
Terminal Columns in Tree Preferences

The terminal columns display...


1. Expected Value - the EV calculated for the active payoff (2).
2. Custom - Node(1) - calculate the EV for a non-active payoff (1).

- 3. Custom - Total_Cost - calculate the value of the variable at the terminal node.
- 4. Path Probability - the path probability for each terminal node of the recommended strategy.
- 5. Custom - PathProb() - the path probability for every terminal node.
- 6. Custom - PathProb()*Total_Cost - the EV multiplied by the path probability

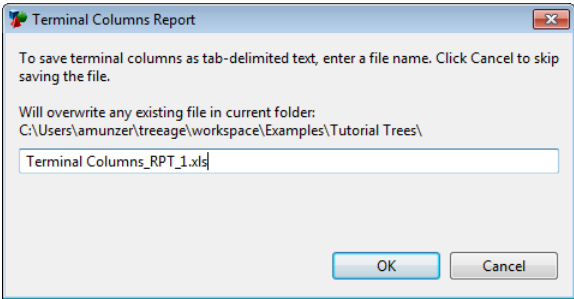
When you roll back the tree, the terminal columns are displayed to the right of each terminal node.



Terminal Columns roll back

 Terminal columns are a convenient way to show extra payoff values at each terminal node. Note that the "Calculate extra payoffs" tree preference must be checked to output additional payoffs.

When you roll back a model with terminal columns, you are given the option to export the terminal column data to a tab-delimited *.xls file in the same folder as the model.



Export terminal column data dialog

The file can then be opened in Excel. If you do not have excel, you can open the tab delimited file in a text editor.

14.4 Other tree display preferences

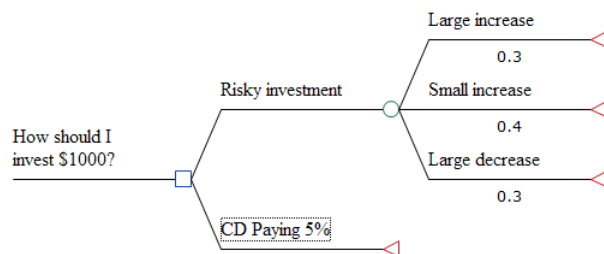
Each tree created in TreeAge Pro can be given its own distinct set of display preferences. Refer to the Tree Preferences Chapter for settings not covered here.

14.4.1 Hiding and boxing payoffs

By default, the active payoff formula is displayed at each terminal node when the tree is not rolled back. This information can be hidden if, for example, you want to simplify a visual presentation of the tree.

To hide terminal node payoff formulas:

- Choose Tree > Tree Preferences from the menu or press the *F11* key to open the Tree Preferences dialog.
- Select the Terminal Nodes preference category.
- Uncheck the option labeled Display payoff names, and press *ENTER* or click *OK*.



Stock tree with payoffs hidden

Or, the tree's payoffs can be displayed in boxes, even when the tree is not rolled back.

To enclose payoffs in boxes when the tree is not rolled back:

- In the Terminal Nodes preferences category, check the option labeled Display payoff names and the option labeled Boxed.

14.4.2 Hiding probabilities and branch labels

The numbers or formulas entered for probabilities can be hidden from view. Node branch labels can also be hidden.

To turn off the display of probabilities and/or node names:

- Choose Tree > Tree Preferences from the menu or press the *F11* key to open the Tree Preferences dialog.
- Select the Node Text/Comments preference category.
- Check the option labeled Hide probabilities only, or the option labeled Hide all node texts. Press *ENTER* or click *OK*.

14.4.3 Displaying a "skeleton" tree

The skeleton tree display option has not yet been implemented.

-
-
-

14.4.4 Terminal node numbers

To show scenario/terminal node numbers:

- Choose Tree > Tree Preferences from the menu or press the *F11* key to open the Tree Preferences dialog.
- Select the Terminal Nodes preference category.
- Check the option labeled Automatic node numbering.
- Enter the Numbering text format. The default entry will simply number the nodes.
- Press *ENTER* or click *OK*.

The text you enter for terminal node numbering must use the caret ("^") as a placeholder for the node number. The caret can be used alone or with additional text, as in "Outcome ^". The terminal node number and text will be displayed whether or not the tree is rolled back.

14.4.5 Number all nodes

Instead of just numbering terminal nodes, TreeAge Pro can apply the numbering format you specify to all nodes in the tree.

To show node numbers for all nodes:

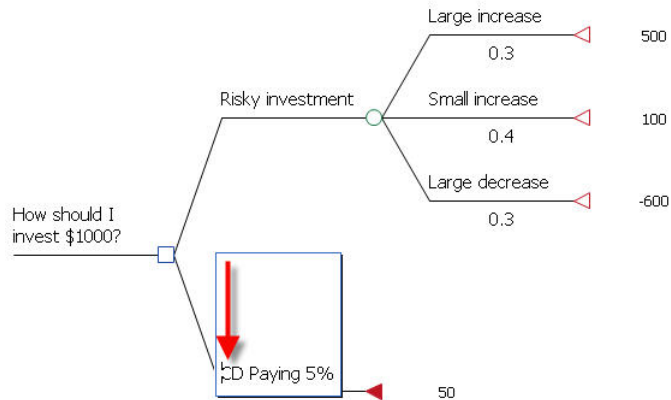
- In the Terminal Nodes preferences, check the option labeled Automatic node numbering. Refer to instructions in prior section.
- Also check the sub-option All nodes in tree.

14.4.6 Increasing or decreasing vertical white space

If two branches that are vertical neighbors appear too close together, there is a way to increase the space between them.

To increase the vertical spacing between two nodes:

- Click on the bottom branch's text label, to the left of the first word in the label.
- Press *CONTROL + ENTER* one or more times, inserting carriage returns until you have created sufficient white space.
- Press *ENTER* to commit the text edits.



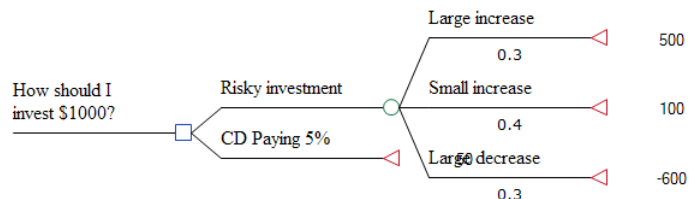
Tree with added vertical whitespace

A tree can be compressed vertically, reducing white space and yielding a very compact tree.

To compress a tree vertically:

- Choose Tree > Tree Preferences from the menu or press the *F11* key to open the Tree Preferences dialog.
- Select the Tree Editing/Layout preference category.
- Check the option labeled Minimize empty space. You must first ensure that Align endnodes is not selected.
- Press *ENTER* or click *OK*.

The figure below illustrates how TreeAge Pro compresses the display of the tree.



Tree with minimized whitespace



This tree preference does not yet account for the position of payoffs. Notice the payoff for the bottom strategy is presented on top of another node.

You can increase vertical white space in conjunction with this tree preference to correct this error manually until this issue is resolved.

If the display of your tree looks broken when the Minimize empty space setting is turned on, review these potential conflicts:

- The Minimize empty space and Align endnodes settings are incompatible.
- Minimize empty space can cause problems when a tree is rolled back, because of a lack of space for roll back boxes.

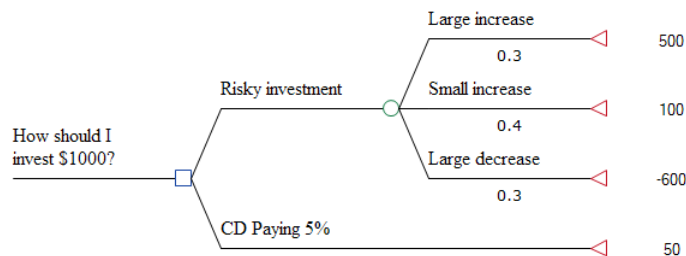
- Using both the Minimize empty space and Branch lines at right angles settings can result in branch lines which slice through node symbols.
- Minimize empty space is likely to cause problems with the display of terminal node columns.

14.4.7 Vertically aligning terminal nodes

There is a quick way to align all terminal nodes in a tree with the right-most terminal node.

To align all terminal nodes at the right edge of the tree:

- Choose Tree > Tree Preferences from the menu or press the *F11* key to open the Tree Preferences dialog.
- Select the Tree Editing/Layout preference category.
- Check the option labeled Align endnodes.
- Press *ENTER* or click *OK*.



Tree with end nodes aligned

For instructions on aligning specific nodes in the tree, refer to the Aligning selected nodes section at the beginning of this chapter.

14.5 Changing fonts

The font, size, and style of any text that appears in a tree can be easily changed. Fonts can either be changed for selected objects or globally, via a tree's preferences.

14.5.1 Changing the font of selected objects

You can change the appearance of text within the model. You can modify the node font for the following attributes: node label, probability, expected value and variables. Each attribute can use a different font.

For notes, the only attribute available is the node text.

To change the font of the selected node:

- Right-click on a single node.
 - Choose Font > [Attribute] from the context menu.
- Each [Attribute] option refers to different text that can be associated with a node.

- Select the font preferences from the Font dialog and click *OK*.



Once a selected node's font has been modified, it is independent of the tree's global font preferences, described in the next section. Changes made to the tree's font preferences will not affect the selected node.

Fonts can be changed for *multiple selected nodes* and/or subtrees in the same manner as above, using the right-click menu on one of the selected nodes.

Fonts can also be changed for notes by right-clicking and choosing Font from the context menu.

14.5.2 Changing font preferences

The Fonts Tree Preferences category makes it possible to specify, for the entire tree, the font used for node names, probabilities, expected value boxes, and (if displayed) definitions of variables. Each button calls up the standard font, size, and style dialog, but changes made in those dialogs apply only in the limited context that their names reflect.

The *Node Font* button will change the branch text font for nodes subsequently created in the tree. It is also applied to existing nodes in the active tree, with the exception of selected nodes and subtrees at which you have individually changed the font.

The *Prob Font* button will change the probabilities font for the active tree. Like the Node Font, this is used for new nodes and existing nodes in the active tree, with the exception of nodes and subtrees at which you have individually changed the probability field font.

The *EV Font* button allows you to change the font for roll back boxes displayed upon roll back of the active tree.

The *Variables Font* button allows you to change the font for variables displayed beneath nodes in the active tree, if you elect to display full variable definitions in the tree.

15. Introduction to Variables and Sensitivity Analysis

This chapter provides a basic tutorial on the use of variables and one-way sensitivity analysis in decision trees.

Subsequent chapters cover TreeAge Pro's many useful tools for working with variables, as well as more complex, multi-way and probabilistic sensitivity analysis. In particular, refer to the first sections of both the Building Formulas Chapter (on defining variables non-numerically) and the More Sensitivity Analysis Tools Chapter (on avoiding sensitivity analysis problems).

Some users may wish to refer to the Index to find out more about special uses of variables in Markov microsimulation, user-defined Python functions, and clones.

15.1 Sensitivity analysis background

Sensitivity analysis was introduced in the Decision Analysis Primer Chapter as a means of assessing the extent to which a model's calculations and recommendations are affected by uncertainty. Specific questions about the model that sensitivity analysis can help answer are:

1. Is a model sensitive to a particular uncertainty — e.g., does varying a parameter's value result in changes in optimal strategy?
2. If a model is sensitive to a particular uncertainty, at what value(s) of the parameter does the model recommend a change in strategy?
3. Does the sensitivity analysis result make sense? (This is a model debugging question.)

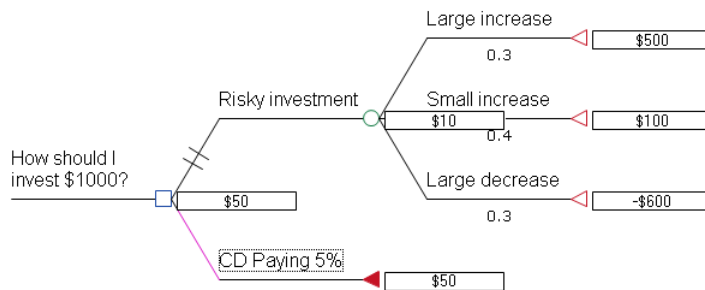
The tutorial in this chapter shows how to prepare a tree for sensitivity analysis, how to perform a one-way sensitivity analysis, and how to interpret the results. The More Sensitivity Analysis Tools Chapter covers multi-way sensitivity analysis and tornado diagrams. The Monte Carlo Simulation Chapter covers the use of probability distributions and Monte Carlo simulation to analyze models with complex or numerous uncertainties.

The Working With Variables Chapter and the Building Formulas Chapter provide important details on working efficiently with variables in decision trees. The information in these chapters can help you improve your productivity when building complex decision trees, and also insure against costly modeling errors.

15.1.1 Variables and sensitivity analysis

Up to this point in the tutorial, decision trees have been analyzed on the basis of baseline, numeric values for payoffs and probability values. In order to perform sensitivity analysis on an uncertain quantity, however, its numeric value must be replaced with a variable — a named parameter.

In the investment decision modeled in the first few chapters, subjective estimates for probabilities and payoffs were used. The resulting tree is shown below, rolled back. The expected value calculations suggest that the CD investment is optimal.



Investment tree rolled back

The basic uncertainty is that, at the time of the decision, the investor cannot know what the price of the equity will be in one year. A simple probability distribution — a chance node with three branches — represents a range of possible changes in the risky investment's value. Assuming that you have a mix of different expert opinions about the risky investment, it would be useful to be able to perform sensitivity analysis on the related parameter uncertainties. The extreme, 10th and 90th percentile values in the payoff distribution — a \$600 decline or \$500 rise in value — are possible candidates for sensitivity analysis. Another option is to vary the probabilities.

The next section of this chapter shows how to replace selected numeric values in the tree with variables. *This is a prerequisite to performing sensitivity analysis in TreeAge Pro.*

15.2 Using variables in a tree

In TreeAge Pro, a variable is a named parameter which functions as a placeholder for a numeric value (or a formula). Variables have a variety of functions in TreeAge Pro, including:

- representing uncertain or unknown values, usually in preparation for sensitivity analysis
- acting as placeholders for mathematical formulas that include functions, tables, spreadsheet links, and other variables (for example in defining a complex payoff)

The tutorial in this chapter focuses on the use of variables as a basis for sensitivity analysis. The Building Formulas Chapter will discuss the creation of payoff formulas using variables.

When building complex models, variables are often used from the outset. The investment decision tree has already been completed using numeric payoffs and probabilities, however, so this tutorial will take a different course, replacing existing numeric values with variables.



Some procedures explained in this chapter are not required steps for using variables in the investment tree. Optional steps will be identified as such.

15.2.1 Steps for using variables

There are three basic steps to remember when using a variable as a parameter in a decision tree:

1. *Declare name* — Based on its intended function in your model, decide on a clear name for the variable (following the naming guidelines outlined below). Add the name to the list of recognized variables in the tree.
2. *Define, assign* — Define the variable at a node, often the root, by assigning it a value (or a formula).
3. *Use* — Anywhere the corresponding value is used in the tree (e.g., payoffs or probabilities), in its place substitute the variable name.

TreeAge Pro offers multiple methods for each step; this chapter illustrates a few possible methods.

15.2.2 Guidelines for naming variables

Variable names must conform to certain rules (similar to Microsoft® Excel's rules for cell names). Each variable name must:

- begin with a letter or underscore character (“_”)
- contain only letters, numbers, and underscore characters
- be no longer than 32 characters

TreeAge Pro will alert you if you try to use an invalid variable name.

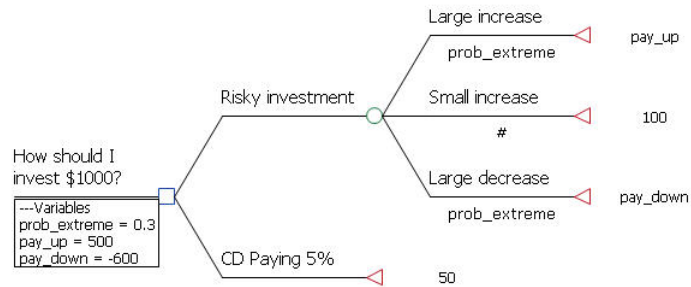
It is recommended that you follow some kind of naming convention when creating variables in a particular model. For example, you could use the prefix “prob” or “p” for probability variables, “c” for cost variables, and so on.

Variable names are not case-sensitive. For example, the names probUp, PROBUP and probup are equivalent; prob_UP would be a different variable, however, since it includes an extra character.

15.2.3 Creating and defining variables

As explained at the beginning of this chapter, there are two apparent sensitivity analysis approaches in the investment tree: vary the extreme probabilities or the extreme payoff values.

The tutorial will start by using variables in the branch probabilities of the extreme outcomes, and after that in the payoffs. The tree will eventually look like the picture shown here.



Stock tree with three variables

It is a good idea, if you already have a working version of your tree without any variables, to keep a backup copy of the tree. For information about creating a project to store your test model, refer to the Managing Projects and Documents Chapter.

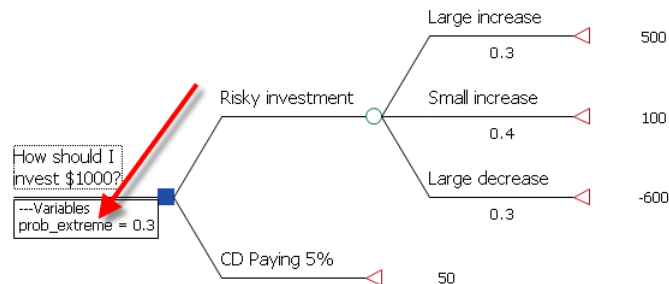
The first step is to create and define a variable that can be used to replace the numeric probabilities of the Large increase and Large decrease branches with a variable placeholder.

To create and define a new variable in a tree:

- Open the Variable Properties View.
- Click the "+" icon in the view toolbar. The Add/Change Variable or Function dialog will open.
- Enter the variable name *prob_extreme* in the name field.
- Enter description and comment information for the variable (optional).
- Enter 0.3 into the Root Definition - Build Expression field.
- Click OK.

Create new variable prob_extreme

The name “prob_extreme” has been added to the list of recognized variable names in the tree, and it has been assigned a default numeric value of 0.3 at the root node. You will see the definition beneath the root node in the Tree Diagram Editor.



Tree showing new variable prob_extreme

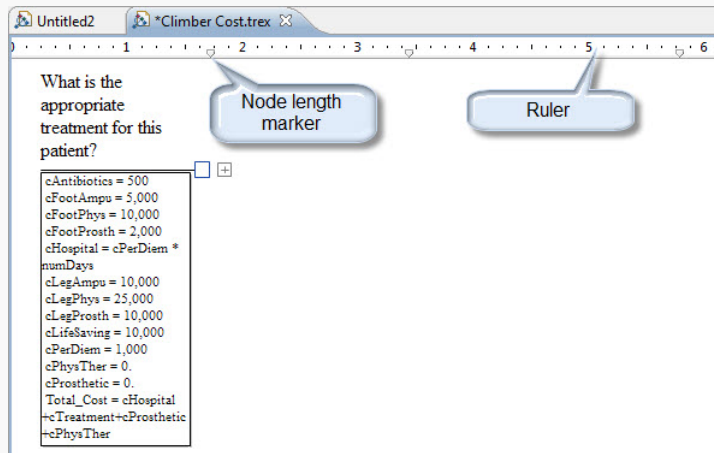
The assignment of a value (or formula) to a variable at a node is called a definition. Which node you define a variable at determines where the variable definition will apply in the tree (i.e., which payoffs and probabilities can use the variable name).

The root node definition of prob_extreme, for example, applies at the root node and *everywhere to the right*, including at the Large increase terminal and at the Large decrease node, both of which will be updated to use the variable.



Use the ruler to stretch a node length to better view variable definitions.

Double-click on the node length marker to stretch a node length to show each variable definitions on a single line.



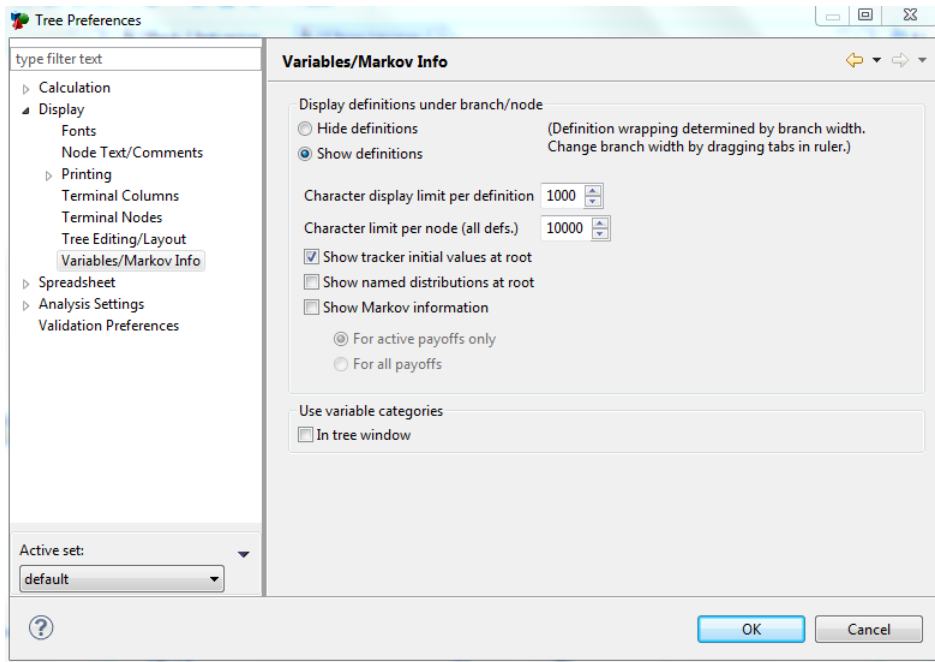
Ruler and node length marker

15.2.4 Finding and fixing problems with definitions

The variable definition, `prob_extreme = 0.3`, should be visible at the root node of the tree. If the definition is not visible below the root node, as in the above picture, it does not mean that the definition does not exist. The first thing to check is whether the display of variable definitions has been turned on in the tree's preferences.

To display variable definitions in the tree:

- Choose Tree > Tree Preferences from the menu or press the *F11* key to open the Tree Preferences dialog.
- Select the category Display > Variables/Markov Info.
- Select "Show definitions" from the "Display definitions at node" options.



Variable display Tree Preferences

If the tree is set to display *variable definitions*, but the definition `prob_extreme=0.3` does not appear at the root node, determine whether the variable was created and defined.

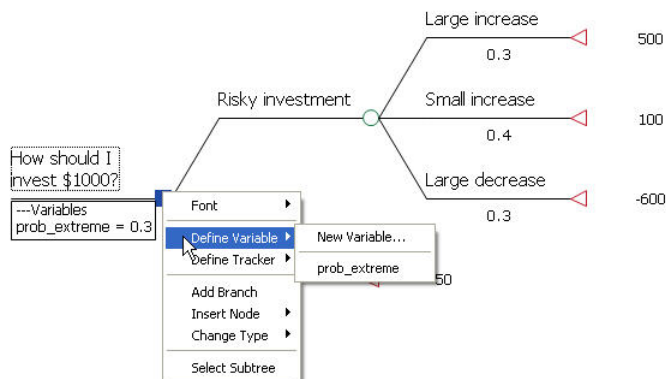


Individual variables can also be hidden/shown in the tree using the Variable Property "Show in tree".

The list of named variables in the tree can be viewed and modified in a number of ways. A quick way to access the variables list is by right-clicking on a node (use the root node for now).

To view the list of tree variables in a "quick" menu:

- Right-click on the root node and choose Define.
- If any variables exist in the tree, their names will be listed (below the New... command).



Variable display in context menu

Later, we will look at using the Variable Properties View and Variable Definitions View to manage a tree's variables.

To open the Variable Properties View:

- Choose Tree > Show View > Variable Properties from the menu.

To open the Variable Definitions View:

- Choose Node > Show View > Variable Definitions from the menu.

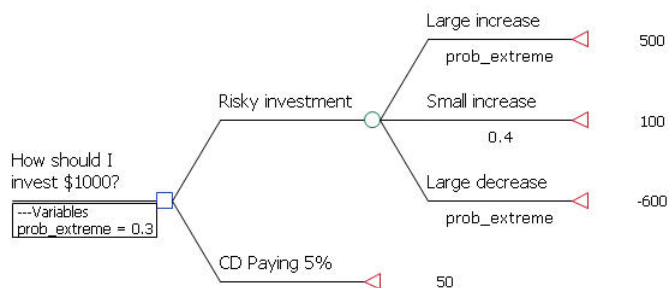
15.2.5 Placing variables in a tree

Previously in this documentation, quantities (payoffs, probabilities, etc.) in the tree were entered as numeric values. However, TreeAge Pro allows for the flexibility of entering quantities as expressions using numbers, variables, tables, functions and distributions. We'll start with the single variable that we created earlier. We'll use two different methods to place the new variable in two probability expressions.

To use the variable `prob_extreme` in the model:

- Select the *Large increase* terminal node and press *SHIFT-TAB* to edit the probability.
- Delete the numeric probability of 0.3, and in its place type a new variable name `prob_extreme`.
- Press *SHIFT-TAB* (or click outside the node) to have TreeAge Pro check the changes you made to the probability expression.
- Click with the mouse in the probability editor, below the *Large decrease* terminal node.
- Click on the elipsis ("...") to open the Formula Editor.
- Clear the 0.3 from the formula editor.
- Select variable in the formula editor's left pane, then double-click the variable `prob_extreme` in the right pane and click OK.

At this point, the variable `prob_extreme` has replaced two numeric value probability values 0.3 in the model.



Stock tree using new variable `prob_extreme`

15.2.6 Adjusting complementary probabilities

Our changes to the probabilities in the tree are not complete. If the value of the variable `prob_extreme` is changed, for example during sensitivity analysis, the probabilities at the chance node will no longer sum to 1.0 — unless a change is made to the 0.4 probability at Small increase.

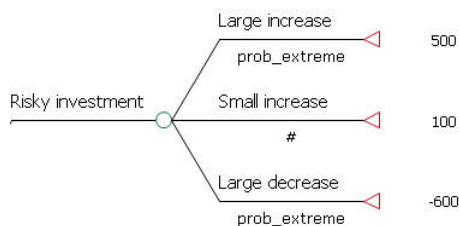
The two options for Small increase's probability are to:

- enter an expression in terms of the new variable (e.g., "1-prob_extreme*2"); or
- use TreeAge Pro's automatic complement calculator ("#").

Let's use the second option, the "#".

To assign a remainder expression to a probability:

- Delete the numeric 0.4 probability of the Small increase node.
- Replace it with # (a hashmark), which will still calculate as 0.4 unless the definition of `prob_extreme` changes.



Use complement calculator with variables

Now, if we change the value of `prob_extreme` to 0.25, the probability of Small increase will automatically be recalculated as $1 - 0.25 - 0.25$, or 0.5, and the chance node's probabilities will continue to be coherent (sum to 1.0).

Once `prob_extreme` has been defined and used properly, and the chance node's probabilities have been adjusted appropriately, you could skip ahead to the section on performing sensitivity analysis, and try analyzing the impact of this uncertainty on your decision.

15.2.7 Create, define and use additional variables

At this point, we have introduced the variable `prob_extreme` into the model. Now we will introduce variables into the payoff expressions for the Large increase and Large decrease terminal nodes.

We will use the same method we used earlier for the first variable and a different method for the second.

To create and define a new variables in a tree:

- Choose Tree > Show View > Variable Properties from the menu.
- Within the Variable Properties View, click the "plus" icon.
- Enter the variable name (`pay_down`), optional description and optional comment.

- Check the box "Define numerically at root" and enter the value -600 into the Value field.
- Click OK.
- Repeat these steps for the variable *pay_up* with value 500.

Now, we will enter the new variables into the payoff expressions for the appropriate terminal nodes.

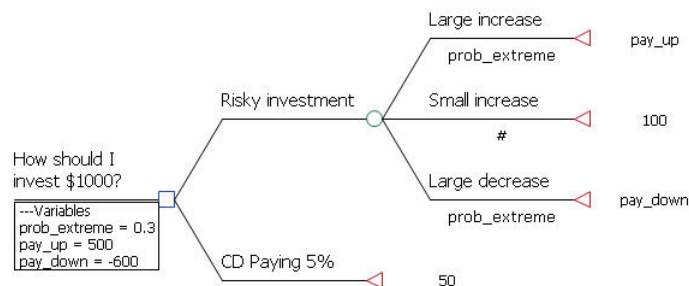
To update the payoff expression for Large increase (Method 1):

- Double-click on the payoff expression 500 to the right of the *Large increase* node.
- In the Edit Payoff dialog, replace the value 500 with the variable name *pay_up* and click OK.

To update the payoff expression for Large decrease (Method 2):

- Right-click on the *Large decrease* node and select Edit Payoffs from the context menu.
- Click on the payoff expression -600 and delete it.
- Click on the elipsis ("...") button to open the formula editor.
- Select variable in the formula editor's left pane, then double-click the variable *pay_down* in the right pane and click OK.

The tree should now look like the figure below - with three variables defined and used in the model.



Stock tree with three variables

The finished version is available as the Get Started tutorial example tree "Three Vars".

15.3 Performing one-way sensitivity analysis

TreeAge Pro can perform sensitivity analysis at a selected node using a range of values for a single variable (one-way sensitivity analysis) or across ranges of values for two or three variables simultaneously (multi-way sensitivity analysis). This chapter covers one-way sensitivity analysis. Multi-way sensitivity analysis, and other advanced sensitivity analysis topics, are described in the More Sensitivity Analysis Chapter.

Start by analyzing the sensitivity of the decision to changes in the probability variable, *prob_extreme*.

To perform a one-way sensitivity analysis:

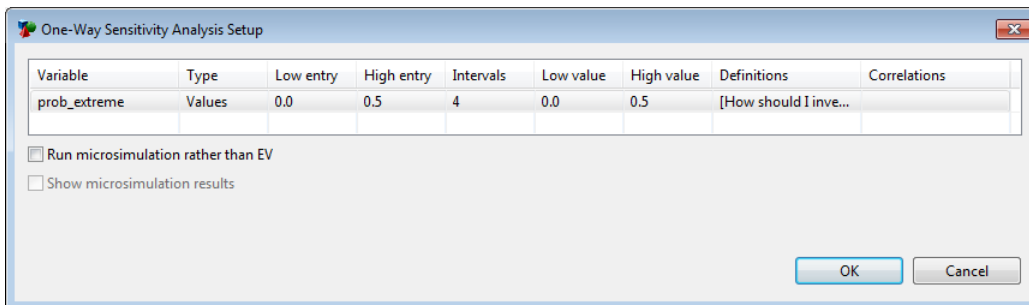
- Open the Three Variables tree you created in the previous section (or open the Get Started tutorial example tree "Three Vars").

- Select the decision node.
- Choose Analysis > Sensitivity Analysis > 1 Way..., or click the toolbar button (see below).
- In the Sensitivity Analysis dialog, click on the Variable column to see a dropdownlist and select the variable prob_extreme.



1-Way Sensitivity Analysis toolbar icon

If you specified a low and high value in the variable's properties, that range will be shown. Otherwise, the range will be set using default values (0 to baseline value). In this case, the saved range is 0 to 0.5.



Sensitivity Analysis Setup Dialog



The checkboxes related to Microsimulation allow you to run a set of trials through the model for each value within the variable range. Refer to the Individual-Level Simulations Chapter for details.

15.3.1 Setting the sensitivity analysis range

The current definition of prob_extreme is 0.3. Recall from earlier in the chapter that our initial interest in sensitivity analysis with the Stock Tree is to test different probability assumptions for the risky investment. For instance, we would like to vary the probability estimates for the extreme outcomes at least from 0.25 to 0.3.

To make the sensitivity analysis more comprehensive, however, we could try the entire possible range for the variable prob_extreme. The tree fragments at left illustrate the concept.

Remember that prob_extreme is used in two branches, with the remainder (or complement) assigned to the middle branch. If we set prob_extreme to 0 (top subtree), the remainder calculated for the Small increase branch will be 1.0. So, prob_extreme=0 can be the minimum value for the sensitivity analysis.

If we try to set prob_extreme to anything above 0.5, probability coherence errors will occur — for instance, at prob_extreme = 0.501 the extreme branches sum to greater than 1.0. The maximum possible value for the variable therefore is 0.5 (bottom subtree). The key estimates for prob_extreme, 0.25 and 0.3, are found within this range.

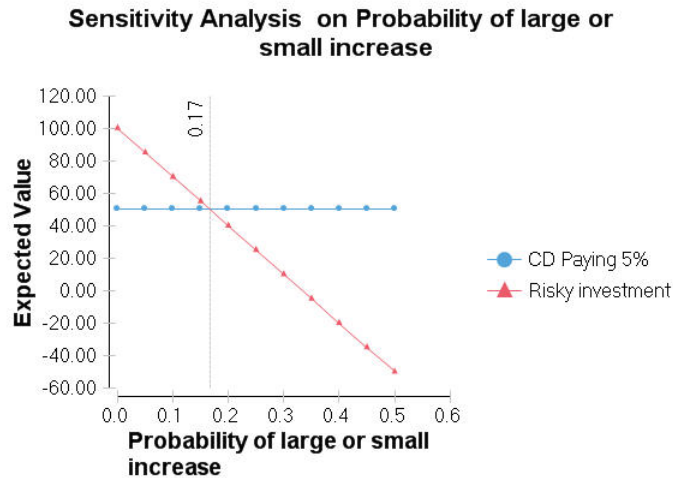
The sensitivity analysis range can be entered in three different ways. Each type is described below with example values applied to the prob_extreme baseline value 0.3.

- *Values*: Enter low and high values for the range directly.
 - Low entry of 0, high entry of 0.5 would be used directly as the sensitivity analysis range values.
- *Adj%*: Enter percentages to be applied to the baseline value to calculate the low and high values for the range.
 - Low entry of 10%, high entry of 20% would be applied to the baseline 0.3 value.
 - Low value would be $0.3 * (1-10\%) = 0.27$.
 - High value would be $0.3 * (1+20\%) = 0.36$.
- *Adj#*: Enter numeric adjustments to be subtracted from and added to the baseline value to calculate the low and high values for the range.
 - Low entry of 0.1, high entry of 0.2 would be applied to the baseline 0.3 value.
 - Low value would be $0.3 - 0.1 = 0.2$.
 - High value would be $0.3 + 0.2 = 0.5$.

Set the sensitivity analysis range:

- Select the Type Values.
- Type 0 for the Low value, and 0.5 for the High value.
- Change the number of intervals to 10. Dividing the range into 10 intervals results in 11 recalculations at the decision node, for prob_extreme equal to 0, 0.05, 0.1, 0.15, 0.2, 0.25, 0.3, 0.35, 0.4, 0.45, and 0.5.
- Press enter or click OK to run the analysis.

TreeAge Pro should immediately begin the analysis. If the analysis were a long one, you could monitor its progress in the status bar at the bottom of the TreeAge Pro window. When it is complete, a graph is displayed.



1-way sensitivity analysis graph

If the analysis does not complete due to errors, read the error message and make note of the node where TreeAge Pro reports a problem. Most likely, the problem is either too wide a range for the uncertain variable, or a problem with the assignment of probability variables in the chance node branches.



If a tree is saved after running sensitivity analysis, the sensitivity analysis range for each variable is saved in the Variable Properties.

15.3.2 The sensitivity analysis graph and report

Because the analysis was done at the decision node, there should be two lines corresponding to the two alternatives, Risky investment and CD paying 5%. Each alternative's expected value is plotted as a function of the increasing value of prob_extreme.

Each strategy's line is composed of line segments connecting the line marker symbols that identify that alternative's expected value at successive intervals of the analysis. A legend to the right identifies the symbol assigned to each particular alternative.

Deviations of a line from the horizontal indicate that strategy's sensitivity to the variable. An alternative represented by a horizontal line in the graph, such as the CD paying 5% option in the example, is unaffected by the changes in the variable. In contrast, the payoff represented by the Risky investment line is an decreasing function of the variable.

The analysis text report, showing the underlying calculated values, can be opened by clicking the Text Report link to the right of the graph.

prob_extreme	Risky investment	CD Paying 5%
0.0	100	50
0.05	85	50
0.1	70	50
0.15	55	50
0.2	40	50
0.25	25	50
0.30	10	50
0.35	-5	50
0.4	-20	50
0.45	-35	50
0.5	-50	50

1-way sensitivity analysis text report

Thresholds are described in the next section.



If the trend of a line in the graph does not make intuitive sense, this may indicate a problem with the definition or use of the variable in the model.

Sensitivity analysis can be used to look also for errors in complex formulas, which might be indicated if changing the value of a parameter (even one that is certain) does not have the anticipated effect on calculations.

The visual elements of the one-way sensitivity analysis graph, like other line graphs, can be customized in a number of ways, as described in the Graphs Chapter. This includes changing graph size, texts, line markers, and numeric formatting.

15.4 Sensitivity analysis thresholds

The sensitivity analysis results can be interpreted graphically.

If two lines in the graph intersect, at the corresponding value of the variable these two alternatives have the same expected value. Crossing points that represent a change in the optimal strategy are called *thresholds*. From the standpoint of expected value, the decision maker should be indifferent between the two options at a variable's threshold value.

15.4.1 Threshold lines

At each threshold, you will see a dotted vertical line. The line stretches up from the x-axis through the crossing point for optimal strategies. The variable value associated with the threshold is shown next to the threshold line.

In the previous section's example, the threshold line marks a crossing point when prob_extreme = 0.17. For values higher than that, including the baseline 0.3 probability, the CD has a higher return and is therefore optimal. For values of prob_extreme less than 0.17, Risky investment is optimal.

In this analysis, the threshold is not close to our two best estimates, 0.25 and 0.3, so perhaps the model is not sensitive to this particular uncertainty.

Among other options for formatting graphs, you could change the numeric formatting of the x-axis to calculate the threshold to a greater degree of accuracy.

15.4.2 Thresholds Report

You can also pull threshold information from the analysis by clicking on the "Thresholds Report" link to the right of the Sensitivity Analysis output graph. This report focuses on the variable value and strategies associated with each threshold in the analysis. See below.

Attribute	Variable	Var. Value	Strategy 1	Strategy 2	Exp. Value
EV	prob_extreme	0.167	Risky Investment	CD Paying 5%	50

Sensitivity Analysis Thresholds Report

Note that the report shows the variable, its threshold values, the strategies that are equivalent at the threshold and the EV value for the strategies at that threshold.

15.4.3 A caveat on thresholds

If all of the lines in a sensitivity analysis graph are straight, the threshold analysis in the graph will be exact. However, if any lines appear curved, the threshold analysis is a linear approximation, and its accuracy will increase as the width of the intervals decreases.

The analysis performs calculations only at the ends of the N number of intervals specified when running the analysis. The lines plotted on the graph are accurate at these N+1 discrete points, but not necessarily in between (e.g., if an alternative's expected value is an exponential or other non-linear function of the variable's value).

The sensitivity analysis graph will not recognize a threshold if the optimal strategy is the same at both ends of the interval, but changes back and forth within the interval. The likelihood of this error can be reduced by increasing the number of intervals, thus reducing their width.



A more sophisticated, non-graphical form of threshold analysis is described in the More Sensitivity Analysis Tools Chapter. This chapter also includes more information on one one-way sensitivity analysis and other kinds of sensitivity analysis.

16. Working With Variables

This chapter expands on the aspects of TreeAge Pro's variables interface introduced in the previous chapter, and also introduces a number of important windows, dialogs, and other tools that facilitate working with variables in trees.

For information on managing tables, refer to the Tables Chapter. Also refer to the Building Formulas Chapter for information on building complex variable definitions and using functions.

Users of the Excel Module can refer to the Using the Excel Module Chapter to learn about features available for managing variables in Excel.

Note that the methods used to manage variable properties and definitions have changed significantly in TreeAge Pro 201x. This chapter will describe the new methods.

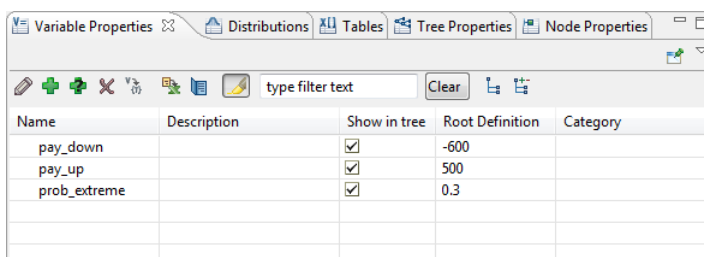
16.1 Variable Properties View

The Variable Properties View is used to manage variable properties. Unlike variable definitions, variable properties apply to the entire tree. Therefore, the Variable Properties View is a tree-level view rather than a node-level view.

To open the Variable Properties View:

- Choose Views > Variable Properties from the toolbar.

Below is an image of the Variable Properties View from the Get Started tutorial example model Three Vars.

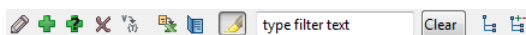


Name	Description	Show in tree	Root Definition	Category
pay_down		<input checked="" type="checkbox"/>	-600	
pay_up		<input checked="" type="checkbox"/>	500	
prob_extreme		<input checked="" type="checkbox"/>	0.3	

Variable Properties View

The main grid contains a list of the tree's variables along with a few of the key variable properties. These properties can be edited within the grid. If you change the variable name, all references to that variable within the tree will be modified as well.

The Variable Properties View toolbar provides additional functions.



Variable Properties View toolbar

The functions associated with the icons are presented below. Each function is described in subsequent sections.

1. Edit Variable
2. Add Variable
3. Add Multiple Variables
4. Delete Variable
5. Convert Variable to Tracker
6. Edit in Excel
7. Generate Variables Report
8. Highlight
9. Filter Text/Clear
10. Show Categories Tree/Group Variables by Category

Variable Properties View toolbar functions

16.1.1 Edit Variable

This function is used to edit the properties of the selected variable. Only one variable can be selected.

The Add/Change Variable or Function Dialog will open with the properties of the selected variable displayed.

The screenshot shows the 'Add/Change Variable or Function' dialog box with the 'General' tab selected. The 'Name' field contains 'pay_down'. The 'Description' and 'Comment' fields are empty. The 'Root definition' section has a 'Build Expression' field containing '-600' and a 'Grid View' button. The 'Add to Expression' section has a 'Group' list with 'Variables', 'Functions', 'Operators', 'Keywords', 'Trackers', 'Tables', and 'Distributions'. The 'Element' field is empty. The 'Calculated value (at How should I invest \$1000?):' field shows '-600.0'. The 'Definition info' field is empty. The 'Show definitions in tree' checkbox is checked. The 'OK' and 'Cancel' buttons are at the bottom.

Add/Change Variable or Function Dialog

Within this dialog, you can edit the variable's properties - name, description, comment. You can also enter a variable definition at the root node either by entering the definition or via the formula editor or content-assist.

Within the Root Definition section, there are two main areas Build Expression and Add to Expression.

1. *Build Expression*: The expression that is being modified by the Formula Editor.
2. *Add to Expression*: The elements to add to the expression are found here and are:
 1. *Group*: Element type to display in the Element section.
 2. *Element*: Individual item to add the the expression.
 3. "+": A Button to add the selected Element to the Build Expression editor. An

When the changes are complete, click OK to save.

Note that the Add/Change Variable or Function dialog contains three property categories.

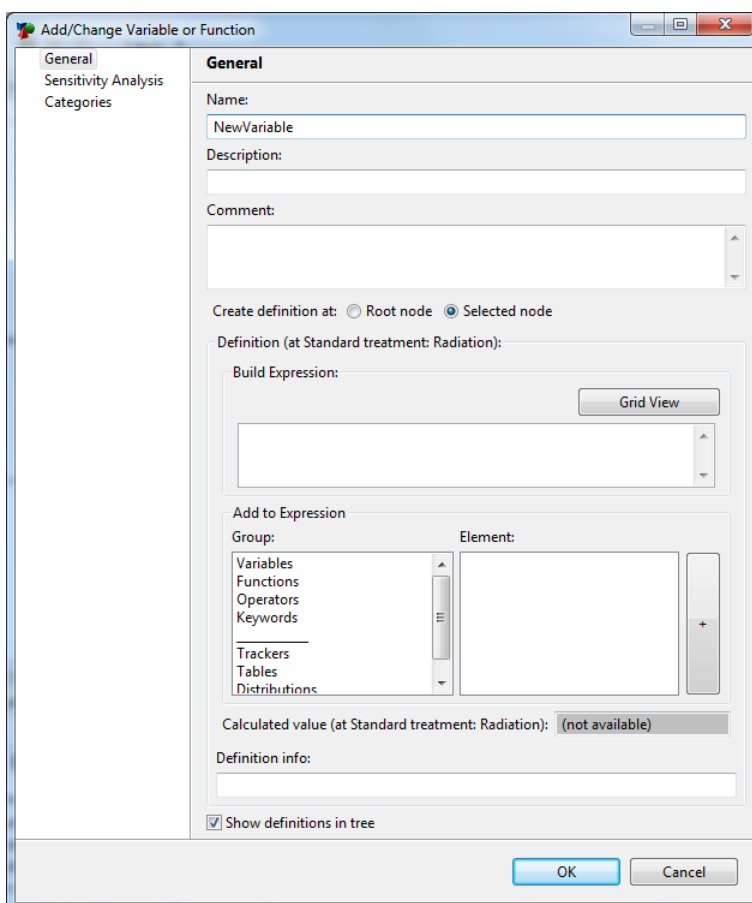
1. *General* - Maintains main variable properties as described above.
2. *Sensitivity Analysis* - Maintains sensitivity analysis range and correlations. Refer to Sensitivity Analysis Variable Properties Section.

3. *Categories* - Maintains Variable Categories associated with the variable. Refer to Variable Categories Section.

16.1.2 Add Variable

This function is used to add a new variable to the model. The Add/Change Variable or Function Dialog (see prior section) will open with a default variable name and default properties (mostly blank). If you have selected to add a variable and you are not at the Root node, you will be able to select where you want to Create the Definition either: *Root Node* or *Selected Node*. A Variable should either be added at the Root node or at the Selected node.

Enter the new variable's name and properties within the dialog.



Add/Change Variable or Function prompting to define at either the Root or the Selected node.

Click OK to save the new variable.

16.1.3 Add Multiple Variables

This function works the same as Add Variable except that you can add a number of variables at a time.

16.1.4 Delete Variable

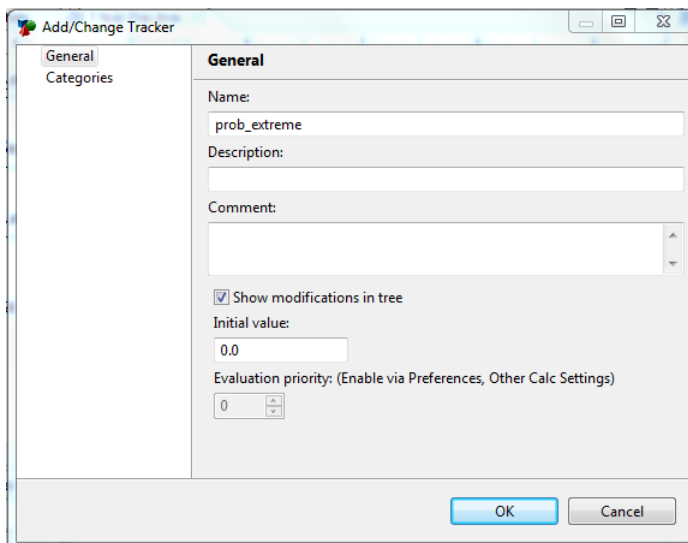
This function is used to delete one or more selected variables from the tree. Deleting variable(s) will also delete all definitions for the affected variable(s).

Expressions that reference deleted variables will no longer function correctly. These expressions will need to be updated.

16.1.5 Convert Variable to Tracker

This function converts the variable into a tracker. Trackers are used in Microsimulation to store and retrieve data associated with individual trials. Refer to the Individual-Level Simulation and Markov Models Chapter for details on the use of trackers.

When executed, this function opens the Add/Change Tracker Dialog to allow you to create the tracker from the existing variable.



Add/Change Tracker Dialog

After you click OK, the variable is converted to a tracker.

16.1.6 Edit in Excel

This function exports the tree's variables to an Excel worksheet. You can then edit the variables within the worksheet and send the new properties/values back to TreeAge Pro.

This function requires the optional Excel Module. It is described in the Using the Excel Module Chapter.

16.1.7 Generate Variables Report

This function creates a report listing...

1. A complete variables list for the the tree including each variable's name and selected properties.
2. For a single selected node, a list of variables defined at or to the left of the node and their properties.

Name	Description	Type	Unused	Formula	Value	DefinedAt	DefinitionInfo	Low	High	Cor	Actions
pay_down				-600	-600	How should I...		0	0		Export Report As...
pay_up				500	500	How should I...		0	0		
prob_extreme				0.3	0.3	How should I...		0	0.5		

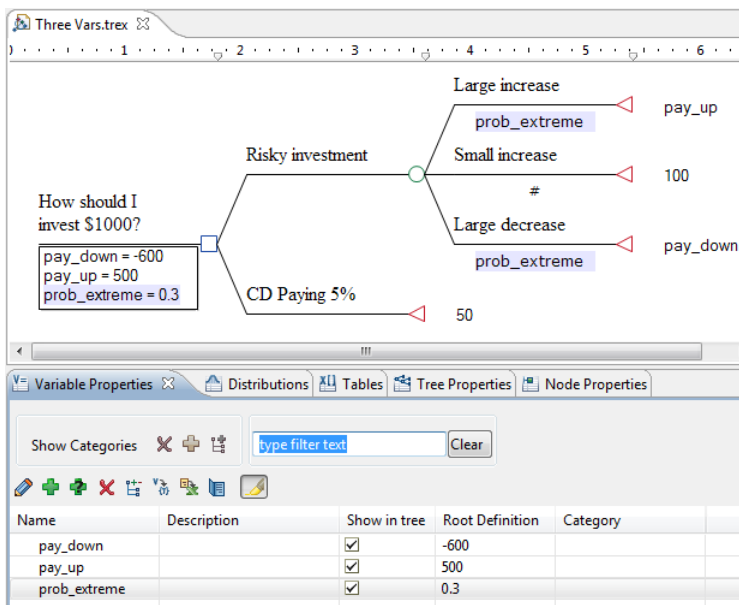
Variables Report

The Variables Report can be useful in determining which variables are unused (and might be deleted). It can also be used to retrieve calculated values based on the applicable variable definition (at the selected node or closest one to the left).

Similar to other reports, the Variables Report can be exported to a number of external formats including HTML and Excel.

16.1.8 Highlight

When this option is depressed, the selected variable is highlighted within the model in the Tree Diagram Editor. See below.



Variable highlighted in model

16.1.9 Filter Text/Clear

The Filter Text field allows you to filter the list of variables to show only variable names that match the Filter Text. For example, if you entered "pay" into the Filter Text, the variables *pay_up* and *pay_down* would be displayed, but not *prob_extreme*.

Enter an asterisk before the expression to look for text in the middle of the variable name rather than the front. For example, if you entered "*down" into the Filter Text, the variable *pay_down* would be displayed, but not *pay_up* or *prob_extreme*.

16.1.10 Show Categories Tree/Group Variables by Category

The Show Categories Tree toggle switch splits the Variable Properties View into two panes, adding a left-side pane for managing Variable Categories. Click the icon again to hide the category pane. Refer to the Variable Categories section for details.

The Group Variables by Category function changes the display within the Variable Properties View to group the variables within the assigned categories.

16.2 Variable Definitions View

The Variable Definitions View is used to manage variable definitions at different nodes within the tree. Variable definitions are created at specific nodes, so the Variable Definitions View is a node-level view. The contents of the view reflect the context of the selected node.

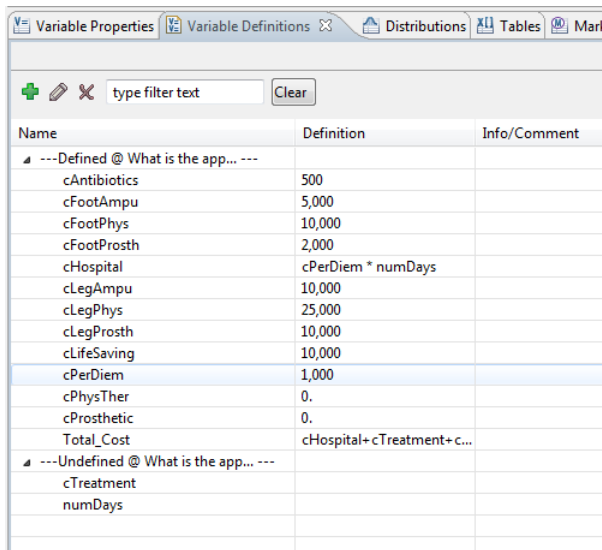


Several Views, such as Variable Definitions View, are node specific. If you have not selected a node or are not on the correct node type, a warning will appear in the View.

To open the Variable Definitions View:

- Choose Views > Variable Definitions from the toolbar.

Below is an image of the Variable Definitions View from the Get Started tutorial example model Climber Cost with the root node selected.



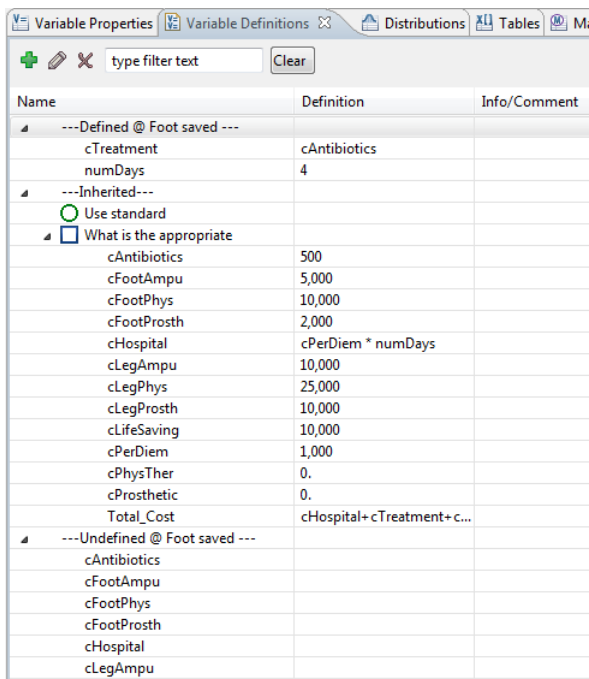
Name	Definition	Info/Comment
---Defined @ What is the app...---		
cAntibiotics	500	
cFootAmpu	5,000	
cFootPhys	10,000	
cFootProsth	2,000	
cHospital	cPerDiem * numDays	
cLegAmpu	10,000	
cLegPhys	25,000	
cLegProsth	10,000	
cLifeSaving	10,000	
cPerDiem	1,000	
cPhysTher	0.	
cProsthetic	0.	
Total_Cost	cHospital+ cTreatment+ c...	
---Undefined @ What is the app...---		
cTreatment		
numDays		

Variable Definitions View

The main grid contains a grouped list of variable definitions. Groups can be collapsed and expanded. Two of the groups are described below.

- *--Defined--* contains variable definitions at the selected node.
- *--Undefined--* contains variables that are not defined at the selected node.

As previously mentioned, the Variable Definitions View is a node-level view. Note how the display changes if we select a different node, specifically via the path *What is appropriate... > Use standard antibiotic > Foot saved*.



Name	Definition	Info/Comment
---Defined @ Foot saved---		
cTreatment	cAntibiotics	
numDays	4	
---Inherited---		
<input type="radio"/> Use standard		
<input type="checkbox"/> What is the appropriate		
cAntibiotics	500	
cFootAmpu	5,000	
cFootPhys	10,000	
cFootProsth	2,000	
cHospital	cPerDiem * numDays	
cLegAmpu	10,000	
cLegPhys	25,000	
cLegProsth	10,000	
cLifeSaving	10,000	
cPerDiem	1,000	
cPhysTher	0.	
cProsthetic	0.	
Total_Cost	cHospital+ cTreatment+ c...	
---Undefined @ Foot saved---		
cAntibiotics		
cFootAmpu		
cFootPhys		
cFootProsth		
cHospital		
cLegAmpu		

Variable Definitions View at the Foot Saved node

At the *Foot Saved* node, only two variables are defined, while others are listed in the -- *Undefined* -- group.

In addition, you can now see the --*Inherited*-- group that contains variable definitions that are "inherited" from nodes to the left of the selected node. The *Foot Saved* node's immediate parent node is *Use standard antibiotic*, which has no variable definitions. Moving one more step to the left, you see the *What is the appropriate...* node (the root node), which has several variable definitions.

The Filter Text field allows you to filter the list of variable definitions to show only variable names that match the Filter Text.

The Variable Definitions View toolbar provides additional functions.



Variable Definitions View toolbar

The functions associated with the icons are presented below. Each function is described in subsequent sections.

1. Add Variable Definition
2. Edit Variable Definition
3. Delete Variable Definition

In addition, a section below describes how to edit variable definitions directly in the view's grid.

16.2.1 Add Variable

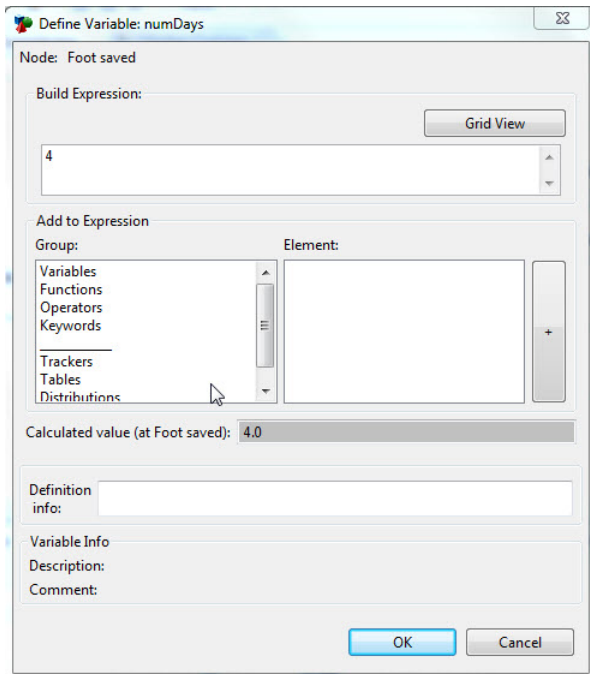
This function is used to add a new variable to the model. This function is the same as the function from the Variable Properties View.

Adding variable definitions is done within the views grid.

16.2.2 Edit Variable Definition

This function is used to edit the selected variable definition. Only one variable definition can be selected.

The Define Variable Dialog will open with the selected variable definition displayed.



Define Variable Dialog

Edit the variable definition in the Build Expression element and click OK to save the changes.

Note that you can select and edit an *--Inherited--* variable definition as well. This will edit the variable definition at the node where the definition currently exists.

16.2.3 Delete Variable Definition

This function is used to delete one or more selected variable definitions from the tree. Deleting variable definition(s) will not delete the variable(s) from the tree.

Only definitions at the currently selected node can be deleted.

To delete variable definitions in the Variable Definitions View:

- Select one or more variable definitions for the currently selected node.
- Click the "X" delete toolbar button.

16.2.4 Edit Variable Definitions in grid

Variable definitions can also be added or edited directly in the Variable Definition View's grid.

For example, if you enter a new definition into one of the *--Undefined--* variables, a new definition is created for that variable at the selected node.

Below is an image of the grid contents before adding a definition.

---	Defined---		
	cTreatment	cAntibiotics	
	numDays	4	
---	Undefined---		
	cAntibiotics		
	cFootAmpu		

Variable Definitions View - Before adding definition

Next, a new definition for the variable cAntibiotics is typed into the grid.

---	Defined---		
	cTreatment	cAntibiotics	
	numDays	4	
---	Undefined---		
	cAntibiotics	501	
	cFootAmpu		

Variable Definitions View - Adding definition

After the new variable definition is added, it can be seen in the *--Defined--* group.

---	Defined---		
	cAntibiotics	501	
	cTreatment	cAntibiotics	
	numDays	4	
---	Undefined---		
	cFootAmpu		

Variable Definitions View - After adding definition

Similarly, you can type a new definition into the *--Defined--* group to replace the existing definition.

16.2.5 Cut/Copy/Paste Variable Definitions

You can cut/paste variable definitions to move them from one node to another. You can also copy/paste variable definitions to copy them from one node to another.

To cut a variable definition in the Variable Definitions View:

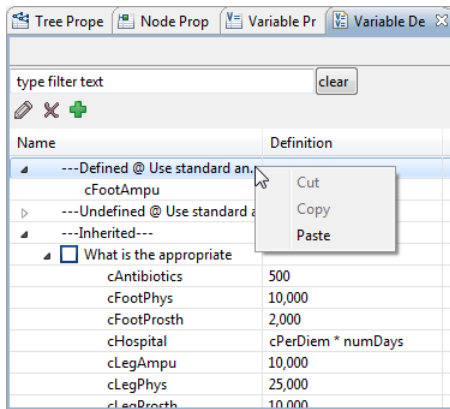
- Right-click on a variable definition.
- Select Cut from the context menu.

To copy a variable definition in the Variable Definitions View:

- Right-click on a variable definition.
- Select Copy from the context menu.

To paste a variable definition:

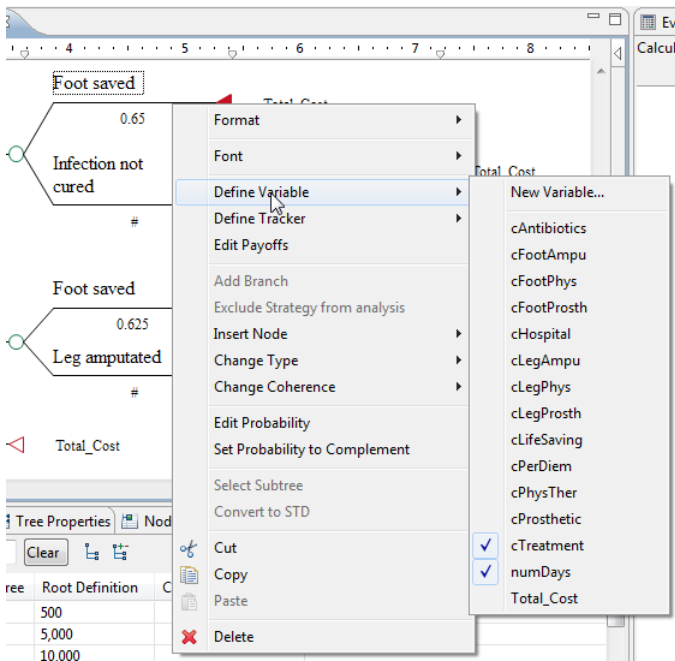
- Select the destination node in the Tree Diagram Editor.
- Right-click on the top row of the Variable Definitions View (refers to variables defined at the selected node).
- Select Paste from the context menu.



Paste Variable Definition

16.3 Defining a Variable at a Specific Node

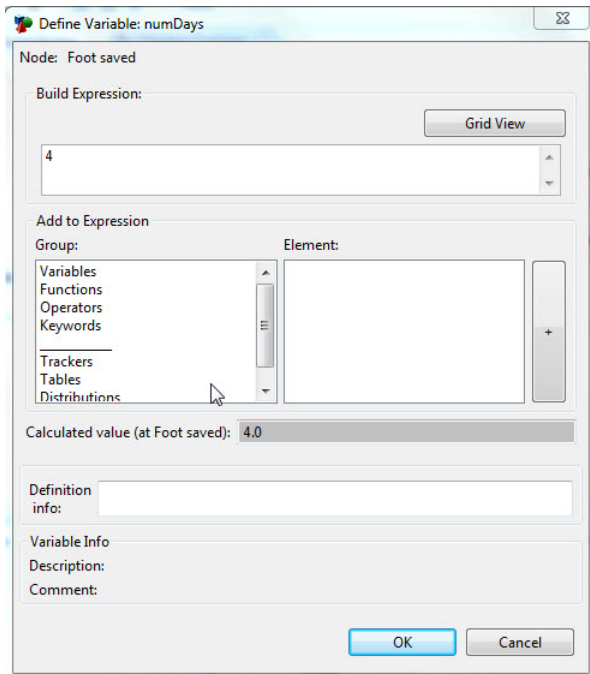
The easiest way to define a new or existing variable at a specific node is to select the node and then right-click. This lets you define a new variable or define an existing variable at the Specific Node from the dropdown menu, as in the figure below.



Defining a variable at the "Foot saved" node by right clicking

In the figure above it indicates both cTreatment and numDays are both defined at the node selected.

To define an existing variable at a specific node, select the variable to define (we use "numDays" in the figure) and the Define Variable Dialog opens. The Define Variable Dialog has been used several times in this and prior chapters. Some additional information on this dialog is provided here.



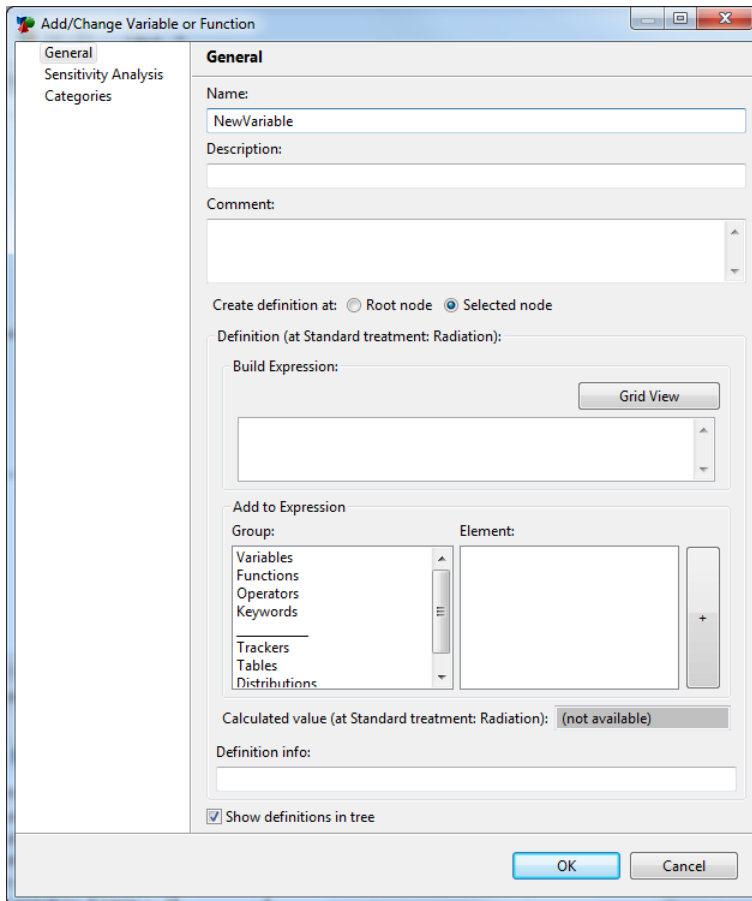
Define Variable Dialog

The Define Variable Dialog is used to add or update a Variable Definition within the context of a specific node. It tells you the Node where you are defining a variable: in this case, *Foot saved*. The read-only Variable Info at the bottom of the dialogue is populated from the the Variable properties. The Build Expression and the Add Expression fields are described in detail in earlier chapters.

The Definition info field is free text that can be used to describe the definition itself.

Another method to *define an existing variable at a specific node* is via the Variable Definitions View. The Variable Definitions View is node specific and the information presented in this view changes depending on which node you have selected in the model. More information of the Variable Definitions View is given in the above section.

To *define a new variable at a specific node*, right-click on the node and choose Define Variable > New Variable... from the dropdown menu. This will open the Add/Change Variable or Function dialog. Note that the "Create definition at" option will default to "Selected node", which you can override to create the definition at the root node.



The Add/Change Variable or Function dialogue for defining a new variable at a specific node with the default variable definition set to "Selected node"

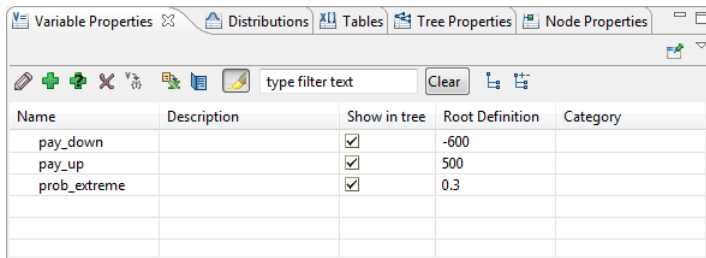
Another method to *define a new variable at a specific node* is via the Variable Definitions View. Once you have selected the node where you want to add a new definition, use the toolbar associated with the Variable Definitions View and select the "+". The Add/Change Variable or Function dialogue will appear. This dialogue will be the same for both methods of adding a new variable, as shown in the figure above.

16.4 Variable Categories

You can use variable categories to help organize long lists of variable definitions in a tree. Use the Variable Properties View to create a hierarchical structure of variables via variable categories.

The categories can be used both in the display of variable definitions in the tree window (under nodes), as well as in the Variable Properties View.

We will use the Get Started tutorial example tree Three Vars as an example. We will categorize the pay_down and pay_up variables under the category "Payoffs" and the prob_extreme variable under the category "Probabilities".

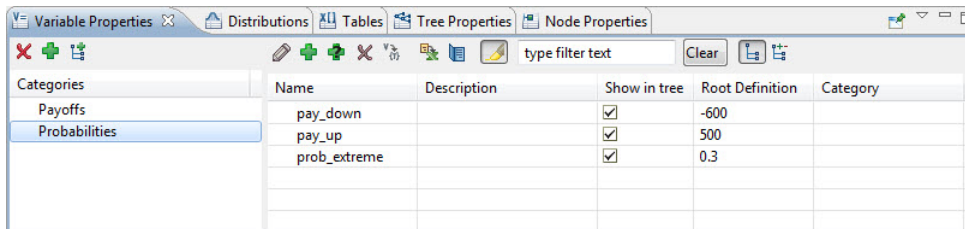


Name	Description	Show in tree	Root Definition	Category
pay_down		<input checked="" type="checkbox"/>	-600	
pay_up		<input checked="" type="checkbox"/>	500	
prob_extreme		<input checked="" type="checkbox"/>	0.3	

Variable Properties View

To create a variable category:

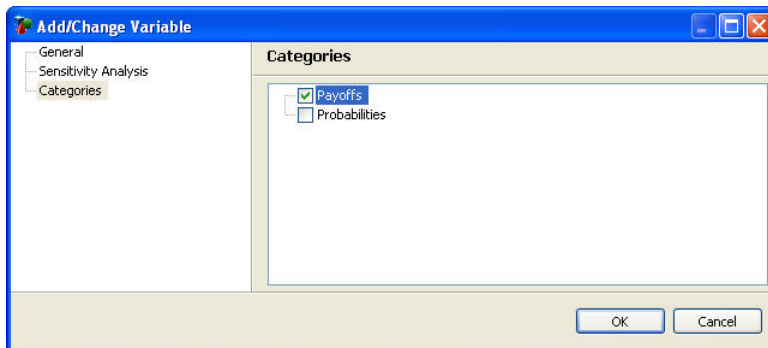
- Click the "Show Categories Tree" button. The categories pane within the view will be displayed to the left of the variables list.
- Click the "plus" icon in the categories pane. A new category "Category1" will be created.
- Click the new category in the categories grid and rename it. In this case, enter "Payoffs".
- Repeat the prior two steps to create the category "Probabilities".



Name	Description	Show in tree	Root Definition	Category
pay_down		<input checked="" type="checkbox"/>	-600	
pay_up		<input checked="" type="checkbox"/>	500	
prob_extreme		<input checked="" type="checkbox"/>	0.3	

Variable Properties View - Categories Pane

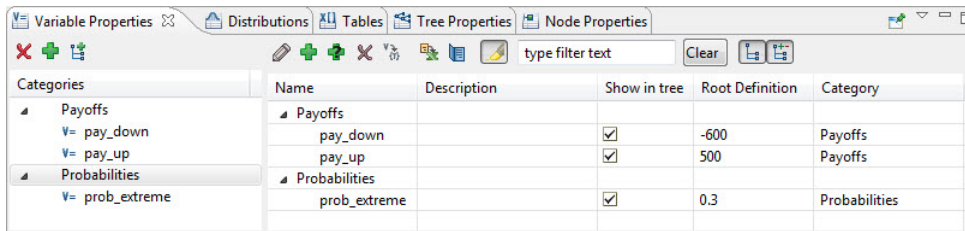
Once the categories have been created, you can assign each variable to one or more categories by dragging the variable onto the category name to the left. You can also assign categories via Categories property category within the the Add/Change Variable Dialog using the Edit Variable function.



Add/Change Variable Dialog - Categories property category

In the figure above, the variable *pay_down* has been assigned to the variable category *Payoffs*.

The Variable Properties View then shows the categories assigned to each variable. Note that the "Group items by category" toggle icon is also depressed.



Variable Properties View - Categories assigned

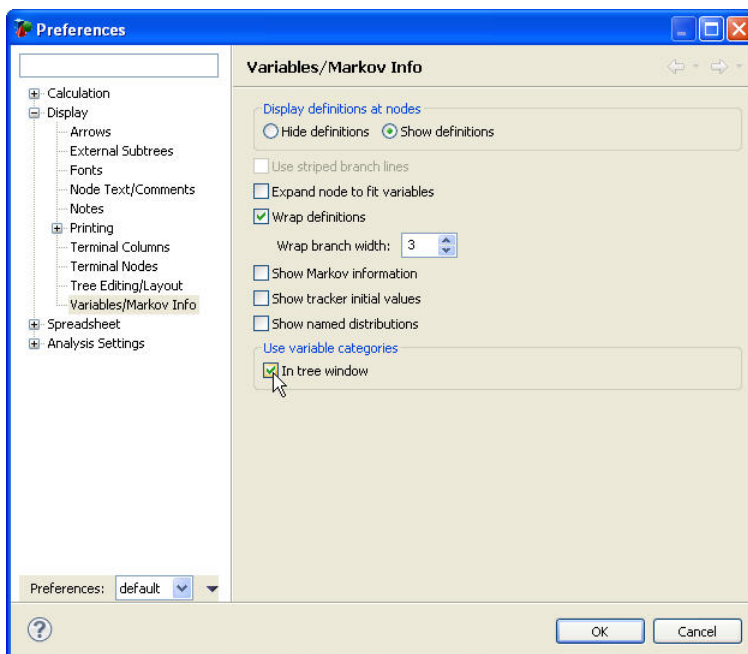
You can create a hierarchical structure of variable categories by creating subcategories within other categories using the "Add sub-category" icon in the categories pane. Variables can be assigned to one or more categories, regardless of each category's position in the hierarchy.

16.4.1 Categories for other inputs

Categories are most-commonly used for variables. However, other model inputs (distributions, trackers, tables) can also be categorized in the same way.

16.4.2 Variable Category Tree Preferences

The tree's variables display preferences allow you to display variable definitions in the model by category.



Tree Preferences - Variable Categories

When checked, the tree shows categories for variable definitions under each node. See below.

The "Low value", "High value" and "Range type" properties are the defaults used when running sensitivity analysis using the specific variable. If the values are changed when running sensitivity analysis, the variable properties are automatically updated to reflect the values used for analysis.

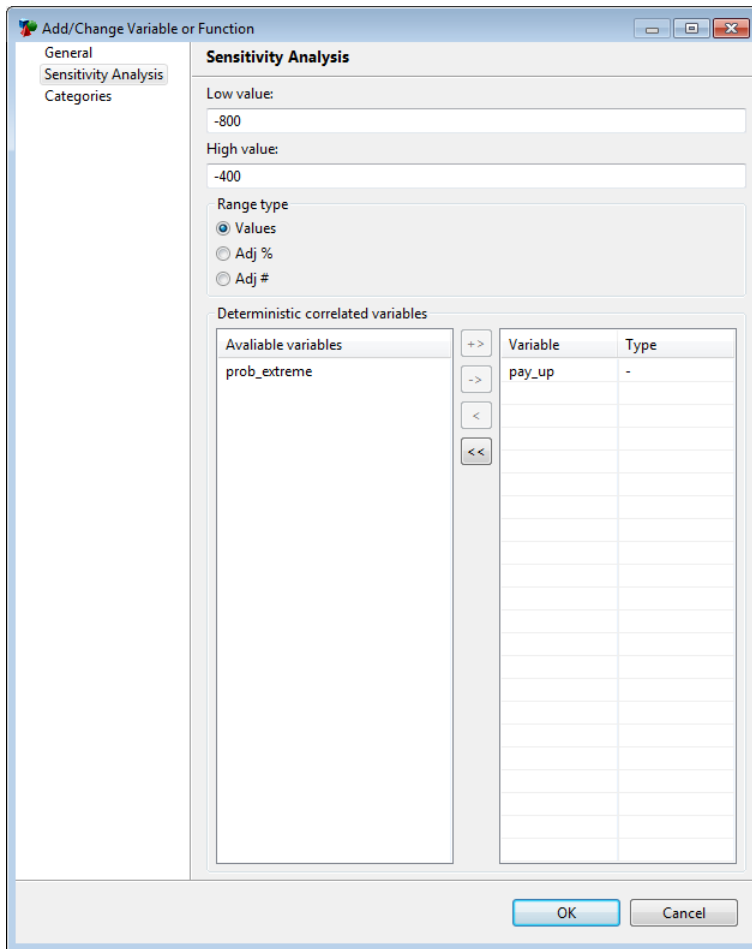
16.6 Sensitivity analysis correlations

Correlations can be set up between any number of pairs of existing variables. When a sensitivity analysis is performed on a variable correlated to another variable, the option is presented to simultaneously vary correlated variables over their own value ranges.

To define variable correlations:

- Open the Add/Change Variable Dialog.
- Choose the Sensitivity Analysis tab.
- Select from the list of available variables.
- Click the "+ >" button for a positive correlation or the "- >" button for a negative correlation.
- Click OK to save the changes.

The linked variables will now be listed to the right of the Correlations buttons, with plus or minus symbols indicating the type of correlation. The identical correlation will show in the properties of the two correlated variables. The correlation can be modified or removed from either variable's Properties dialog.



Add/Change Variable Dialog - Correlation

To remove a correlation:

- Open the Add/Change Variable Dialog.
- Choose the Sensitivity Analysis tab.
- Select from the list of correlated variables (the list to the right).
- Click the "<" button.
- Click OK to save the changes.

You can also click the "<<" button to remove all correlations. To change a correlation's type (e.g., from negative to positive), you must remove the existing correlation and recreate it with the proper correlation.

Sensitivity analysis using variable correlations is described in the More Sensitivity Analysis Tools Chapter.

16.7 Variables testing tools

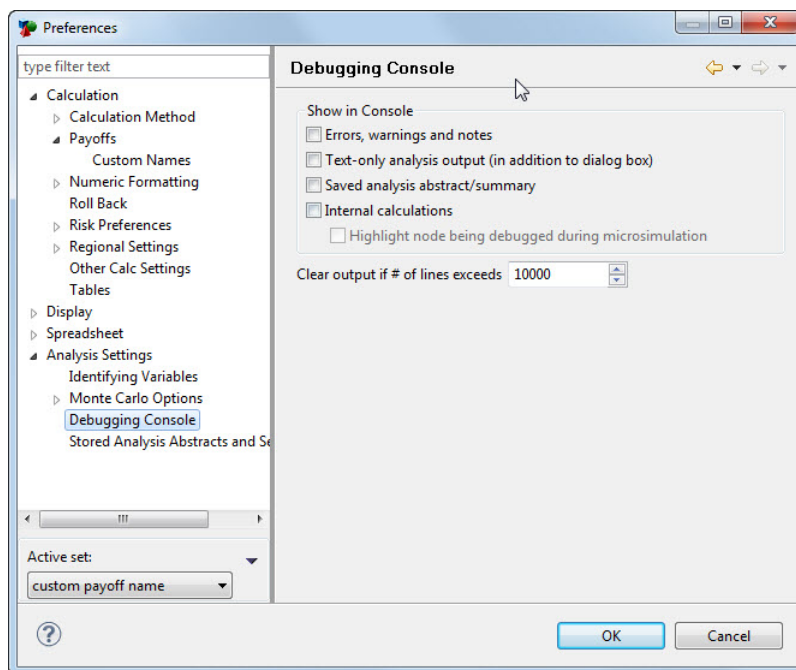
In addition to using TreeAge Pro's Analysis menu commands to see the results of node calculations, TreeAge Pro also provides tools for testing individual variables.

16.7.1 Debugging variable calculations during analysis

The Calculation Trace Console can be used to report on every variable calculation during an analysis. This feature can be used to search for problems in complex formulas.

To turn on detailed calculation debugging:

- Choose Tree > Tree Preferences from the menu or click the *F11* key.
- Select the preferences category Analysis Settings > Debugging Console.
- Check the "Internal calculations" box.



Tree Preferences - Debugging category

Reporting all variable calculations will slow down calculations, so this setting should be turned off except when debugging.

Refer to the Tools and Functions for Complex Trees Chapter for more tips on debugging preferences.

16.7.2 The Evaluator View

The Evaluator View is designed for testing variables and formulas - like a calculator. Any valid expression can be entered into the Evaluator View. The expression is calculated *within the context of the currently selected node*.



This Calculator/Evaluator is an Expected Value-based tool. Distributions return their mean values, and Markov counters and trackers return their default values. For calculations within the context of a Markov Cohort Analysis or Monte Carlo simulation, debugging output may be more useful.

To calculate a variable or formula at the selected node:

- Select the node at which you want to perform the calculation.
- Click on the "=" icon in the application toolbar to open and/or activate the Evaluator View.
- Enter the expression you want to calculate into the top pane of the Evaluator View.
- Click on the "Calculator" icon in the Evaluator View. The calculated value will appear in the bottom pane of the Evaluator View.



Calculator icon

In the following model Climber Cost, from the Get Started tutorial examples, the variable Total_cost is calculated at the selected node, Amputate foot. Note that the calculated value is \$24,000.

The screenshot shows the TreeAge Pro software interface. The main window displays a decision tree with the following structure:

- Root node: **Amputate foot** (Terminal node, Total_Cost)
- Branch 1: **Use experimental antibiotic** (Chance node, 0.625)
 - Branch 1.1: **Foot saved** (Terminal node, Total_Cost)
 - Branch 1.2: **Leg amputated** (Terminal node, Total_Cost)

The **Amputate foot** node is selected, and the **Evaluator** view is active. The formula editor shows the expression `Total_Cost`. The result is **24000.0**. The **Calculation Trace Console** at the bottom shows the expression at the node and the values of various variables:

```

Expression at node: Amputate foot / Amputate foot / thread 0 / _sample 1:
Total_Cost
cPerDiem = 1000.0
numDays = 7.0
cHospital = 7000.0
cFootAmpu = 5000.0
cTreatment = 5000.0
cFootProsth = 2000.0
cProsthetic = 2000.0
cFootPhys = 10000.0
cPhysTher = 10000.0
Total_Cost = 24000.0

Total_Cost = 24,000
  
```

Evaluator view in action

Tree Preferences were set to show internal calculations, so the Calculation Trace Console shows the inputs that contributed to the value of `Total_cost`.



Note that the Evaluator View includes formula editor lookup frames to the right and also supports auto-fill.

16.8 Formula Editor

The Formula Editor in TreeAge Pro helps you create expressions consisting of variables, functions, distributions and all other quantity values in a tree. The formula editor specifically includes the option to select variables to include in your expression.

The Formula Editor is described in detail in the Building Formulas Using Variables and Functions Chapter.

16.9 Variable Definition Arrays

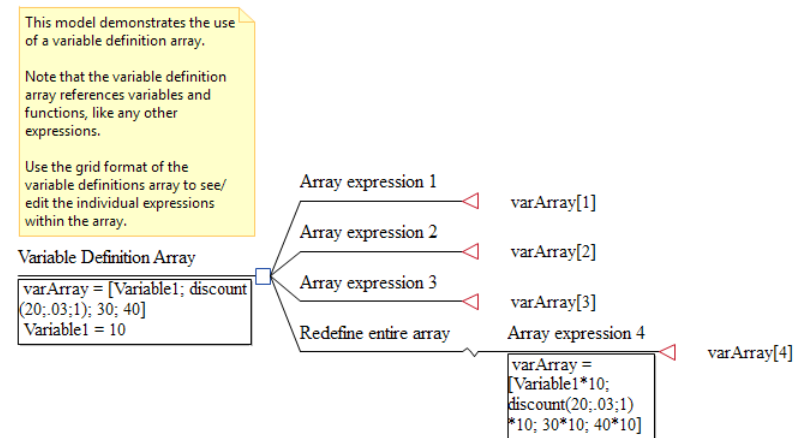
Variable definitions can consist of an array of expressions that can be referenced by index. For example, you could create a variable definition like this...

myVarArray = [10; 20; 30; 40]

Then any reference to the variable *myVarArray* would require an integer index value between 1 and 4, to reference one of the four expressions in the array. For example, *myVarArray*[3] would return the value 30.

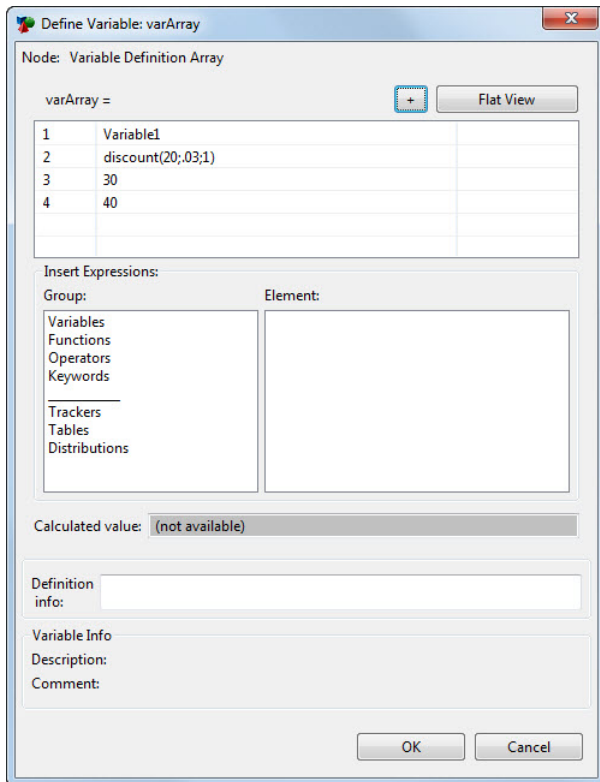
The individual expressions within the array can reference any other variables, trackers, tables, distributions, functions, etc. However, the array cannot refer recursively to itself.

The Special Features tutorial example model, Variable Definition Array, demonstrates this technique.



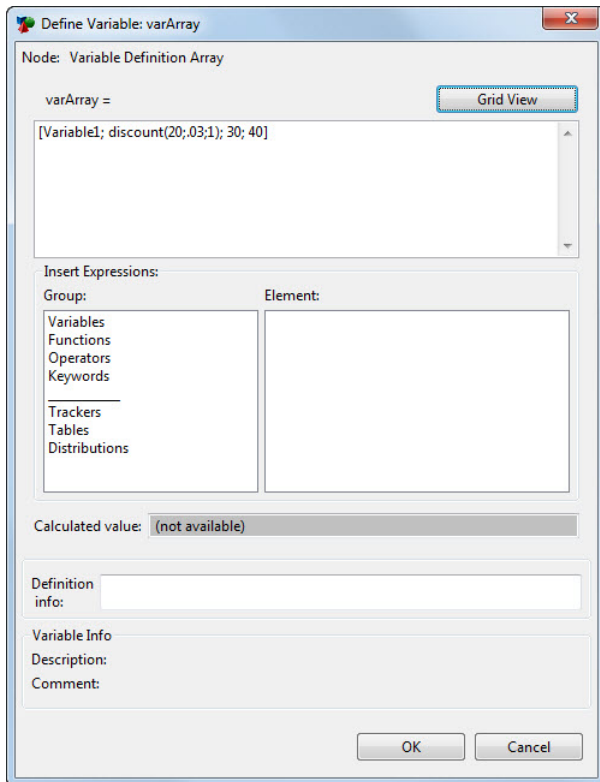
Variable Definition Array Model

Note that when you look at the root node definition for *varArray* in the Define Variable dialog, it is automatically presented in grid format.



Define Variable Dialog in Grid mode

The "+" button adds another row to the grid. The "Flat View" button presents the variable definition as an array with the individual expressions presented together within square brackets and separated by semicolons.



Define Variable Dialog in Flat mode

Use the Flat mode to eliminate rows by deleting an expression and a semicolon.

16.9.1 Using Variable Definition Arrays with Tables

Suppose you wanted to reference different complex expressions within a table. This is not possible with a table alone because tables support only numeric expressions. A variable definition array alone might not be sufficient if you wanted to be able to associate the expressions with specific lookup index values as is done with tables. However, a combination of tables and variable definition arrays provides a possible solution.

For example, let's say you wanted to use different expressions in a Markov model for different age ranges as presented below.

Age	Expression
0-10	cEarly * rr1
11-20	cTeen * rr2
21-40	cAdult * rr3
41-60	cOldest * rr4

Expressions for variable array

You could first setup a variable definition array to cover all the individual expressions. The array is shown below in Flat view.

```
myVarArray = [cEarly * rr1; cTeen * rr2; cAdult * rr3; cOldest * rr4]
```

You could then setup a table that returns the proper variable array index for each age range (using interpolation).

Index	Value
0	1
10	1
11	2
20	2
21	3
40	3
41	4
60	4

myTable table data

The expression would then use the table to determine the proper expression within the variable array as follows.

```
myVarArray[ myTable[age] ]
```

17. Building Formulas Using Variables and Functions

This chapter focuses on how to use variables to build complex formulas for any expression in the model, including payoffs, variable definitions, probabilities, etc.

The Introduction to Variables and Sensitivity Analysis Chapter covers using variables for the purposes of sensitivity analysis on parameter uncertainties in the model.

17.1 Quantity expressions in a model

Quantities must be included in a model in order to analyze the model. Some of the quantities in models are:

1. Payoffs/rewards
2. Probabilities
3. Variable definitions
4. Distribution parameters
5. Function arguments

Quantities in models

Although these quantities can be entered for as simple numbers, TreeAge Pro allows the entry of numeric expressions, consisting of any or all of the following elements.

1. Numbers
2. Variables
3. Table lookups
4. Distributions
5. Functions
6. Operators
7. Keywords

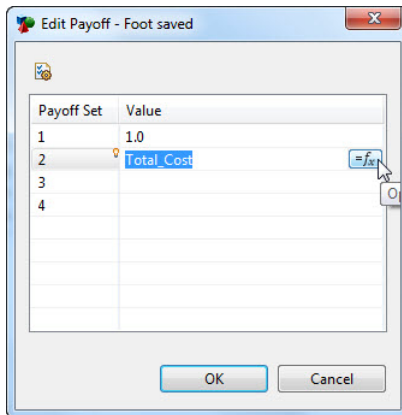
Elements of expressions

There are different reasons to use specific elements within an expression. This chapter does not discuss the reasons. Rather, this chapter focuses on how to construct simple and complex expressions.

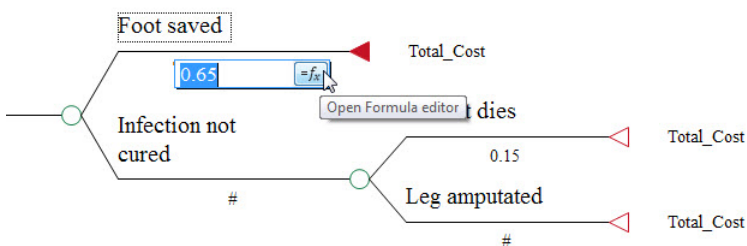
17.2 Formula Editor

The Formula Editor is a tool to create expressions from expression elements.

Within TreeAge Pro, there are many inputs where an expression is required. In most of those inputs, you can open the the Formula Editor to help build the expression. You open the Formula editor by clicking on the "=fx" button within the expression entry. Below are a few examples using the Get Started tutorial example model Climber Cost.



Open Formula Editor from Payoff entry dialog

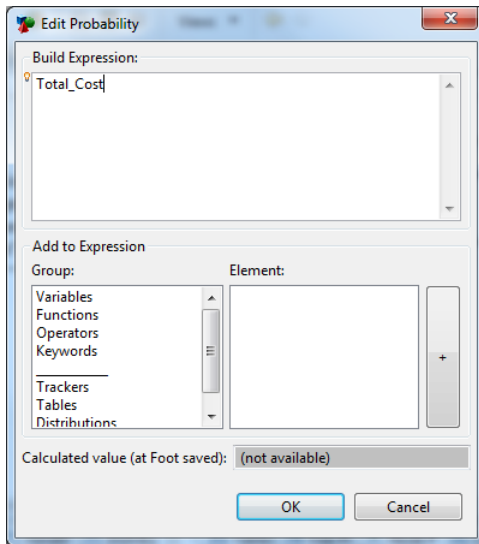


Open Formula Editor from Probability entry

Once the proper expression has been built using the Formula Editor, the expression is placed back in the tree based the original expression input from which the Formula Editor was opened.

The Formula Editor itself is split into two sections which have different elements in them.

1. *Build Expression*: The expression that is being modified by the Formula Editor.
2. *Add to Expression*: The elements to add to the expression are found here and are:
 1. *Group*: Element type to display in the Element section.
 2. *Element*: Individual item to add to the expression.
 3. *"+"*: A button to add the selected Element to the Build Expression editor. An alternative is to double-click the left mouse key.



Formula Editor: The window tells you what you are editing, in this case "Edit Probability"

If you click on an item in the Group section, the Element section will show elements that match the selected type.

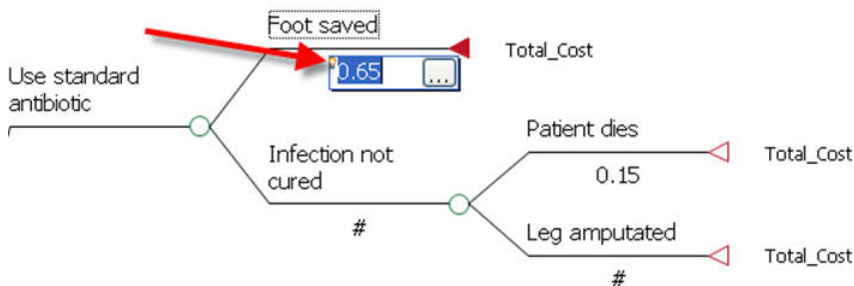
If you double-click on an item in the Element section, that item is added to the expression. Alternatively you can use the "+" button.

Click OK to close the Formula Editor and replace the original expression in the tree with the new expression from the Formula Editor.

Click Cancel to close the Formula Editor and leave the original expression in the tree unchanged.

17.3 Content-Assist

Many expressions in TreeAge Pro allow you to use the content-assist feature. Content-assist is available whenever you see the small light bulb to the left of the expression field. The figure below shows the content-assist indicator in a probability expression field within the Get Started tutorial example tree, Climber Cost.



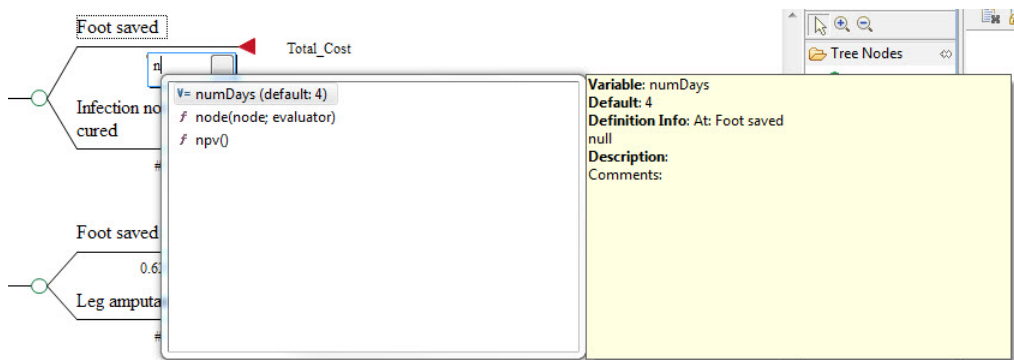
Content-assist option in probability expression

Content-assist attempts to help you complete any partially-typed element in the expression.

To trigger content-assist:

- Select a field with content-assist enabled.
- Begin typing an element.
- Press *Control + Space* on the keyboard.

For example, if you click on the probability expression in the figure above, type the letter "n", then trigger content-assist, you are presented with options to complete the element starting with "n". See below.



Using content-assist in probability expression

Note that you are presented with several possible elements starting with "n" - the *numDays* variable from the tree and the *node* and *npv* functions built into the software. A brief description of each available item is presented in the off-white panel. In the figure above, you can see that the variable *numDays* has the value 4 at this node. If you select one of the content-assist options, it becomes part of the expression.

If no text is entered prior to triggering content-assist, then all possible options are presented.

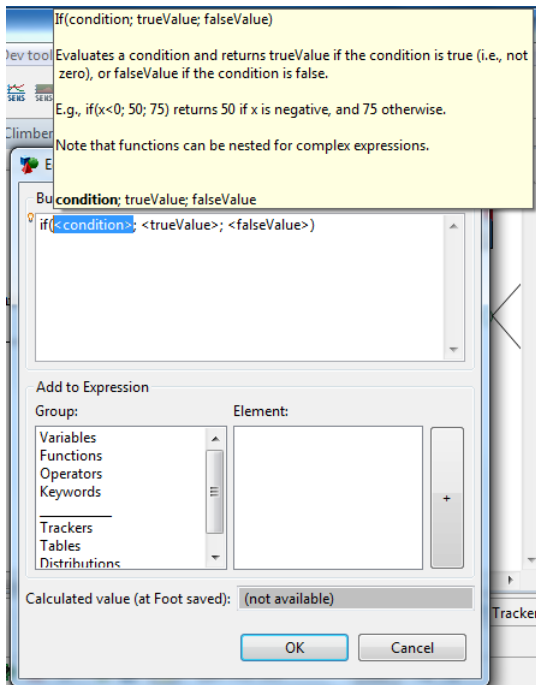


Content-assist works with any element within the expression. It is not limited to the expression as a whole.

17.4 Function Helper

The Function Helper attempts to provide guidance on how to use built-in functions. The Function Helper is available within the Formula Editor and in most expression inputs.

In the Formula Editor, when a function is added to the expression via double-click, micro-help related to that function is immediately presented. In addition, placeholders are added as function arguments. The first argument placeholder text is selected, so you can immediately replace it with the appropriate data/expression for that argument.



Function Helper from Formula Editor

After entering the first argument, press the *tab* key or enter a *semi-colon* argument separator, and the next argument placeholder is selected for entry. This continues until all arguments are entered. *Shift-tab* will return to the prior argument.

If you choose a function via content-assist, the same Function Helper micro-help will appear with the argument placeholders.

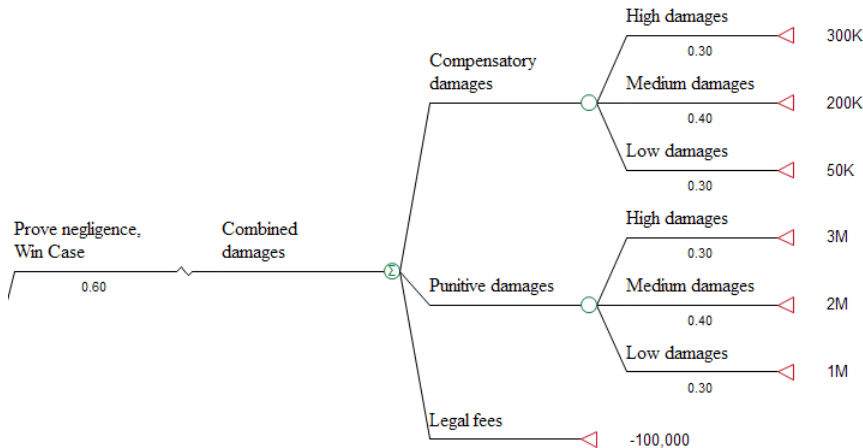
If you type a function into an input field or into the formula editor, the same Function Helper micro-help will appear, but without the argument placeholders, after you enter the open parenthesis after the function name.

17.5 Summation nodes

A summation node can be used to add together the values from portions of a tree. During calculations, the expected value (EV) of a summation node is equal to the sum of the EVs of all branches.

A chance node's EV is equal to the weighted average of its branches EVs. The summation node functions similarly except that the "weight" of each branch would be 100%.

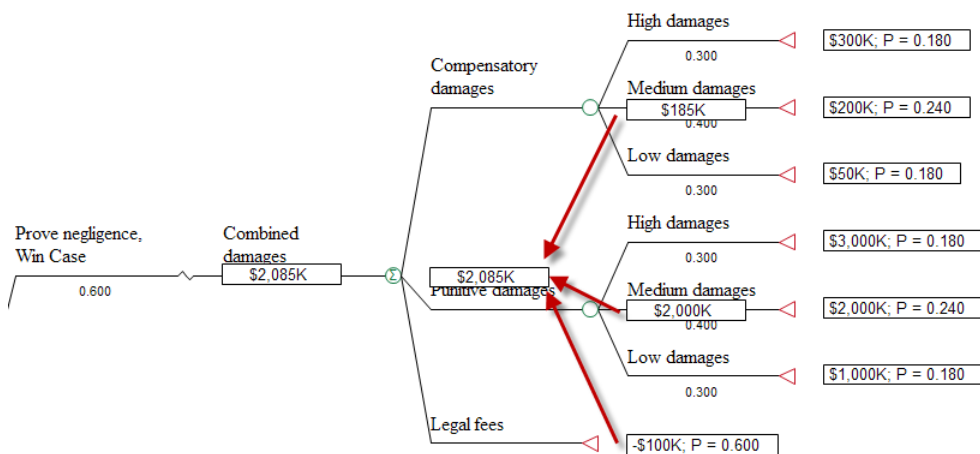
The Legal tutorial example tree, LegalTreeSummation, illustrates the use of the summation node.



Legal tree with summation node

Note that the EV of the legal case if negligence is proven is the combined EV of compensatory damages and punitive damages, less legal fees. The summation node allows you to present the three value elements separately and explicitly, while combining them for the total value of winning in court.

When the tree is rolled back, you can see that the EV at the Combined damages node is equal to the sum of the EVs for its three branches.



Legal tree with summation node - rolled back



Note that summation nodes cannot be placed to the right of a Markov node.

17.6 User-defined python functions

User-defined functions can be created within variable definitions, using the Python script language.

User-defined functions are entered as variable definitions, using a regular variable and the Define Variable Dialog. Python functions require indenting and carriage returns, so the Define Variable Dialog window's behavior changes slightly when it recognizes you are entering/editing a user-defined function.

Refer to the Tools and Functions for Complex Trees Chapter for more details.

17.7 Variable formula examples

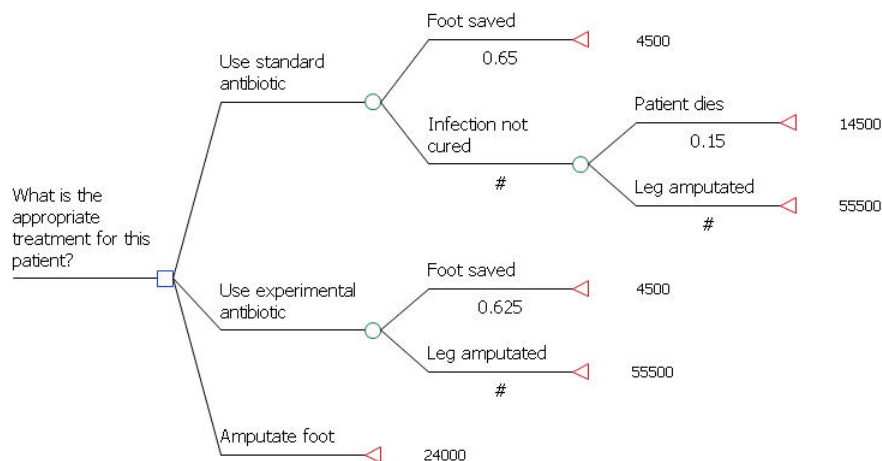
This chapter will continue the tutorial on variables by explaining, in detail, the logic that dictates where variables should be defined in a tree and how variables are evaluated during tree calculations. Provided a basic understanding of how TreeAge Pro searches for variable definitions, it will become easier for you to identify the best location for each variable definition you create.

17.7.1 Building a complex cost formula

The example model used in this chapter is based on the following medical treatment scenario. The assumption is that you are evaluating a new pharmaceutical treatment used in your hospital for advanced infections in an diabetic population. Initially, the modeling of the new intervention is being done on a cost basis, looking only at foot infections.

Previously, the treatment decision was between prompt amputation of the foot, and a course of high-dose, intravenous antibiotics. The new intervention is an antibiotic that works much faster than the old drug, so you will know sooner if it is going to halt the infection and save the foot; soon enough to have complete certainty that performing an amputation below the knee will avoid any mortality risk. For this reason, the new drug has already been adopted.

However, the new drug actually halts fewer infections over its shorter course of treatment, which results in a higher number of amputations. Based on the experience of your medical center, you estimate the costs and probabilities and put them into a decision tree, shown below.



Cost Formula tree

The figure above is of the Special Features tutorial example tree, Cost Formula. As shown in the model, the original high-dose, intravenous antibiotics combined with surgical debridement (removal of tissue) offers a 65% probability of curing the infection and saving the foot. If the antibiotics do not

stop the infection, there is an 15% probability of death with the remaining 85% surviving and requiring amputation at the knee.

The new, experimental antibiotic has a different set of outcomes. It has a 62.5% probability of curing the infection, with the remainder having amputation at the knee.

With the immediate amputation of the foot option, the outcome is assumed to be certain: the patient will survive. With immediate amputation of the foot, or later amputation at the knee, a prosthesis will be fitted and physical therapy will be required.

The following table contains the component costs that are used to calculate total cost for each scenario.

Parameter	Value
Regular antibiotic	\$500
New antibiotic	\$500
Inpatient cost per day*	\$1,000
Foot amputation	\$5,000
Foot prosthesis	\$2,000
Loss of foot, physical therapy	\$10,000
Leg amputation	\$10,000
Leg prosthesis	\$10,000
Loss of leg, physical therapy	\$25,000
Life saving measures	\$10,000

Cost formula parameters



* To calculate the basic cost of the hospital stay associated with the various scenarios, the number of inpatient days for each is also estimated and multiplied by the per diem cost.

The numeric cost payoffs specified for each outcome in the Cost Formula tree have been hand-calculated using a formula combining the appropriate costs for each particular scenario, including costs of hospitalization, drugs, surgery, prosthetics, physical therapy, and other care.

As reflected in the rolled back tree, shown on a previous page, the least costly option is to treat with the standard antibiotic. A Ranking analysis can be used to calculate the additional cost per patient of the experimental antibiotic, \$3427 (or approximately \$70000 for each death that is averted).

17.7.2 Implementing a cost formula using variables

The baseline information may be useful in a budget impact calculations. However, you may want to analyze the model on the basis of different estimates of component costs, some of which are uncertain.

To do this kind of analysis manually would be tedious, so you decide to implement the cost calculations using a cost formula and perform sensitivity analysis on the component variables.

Start by making a copy of the tree:

- Open the Special Features tutorial example tree, Cost Formula.
- Save a copy of the tree, changing the name to *Cost Variables*, and save it in a separate project.

To save some time, a list of definitions of the cost component variables has already been pasted into the tree, at the root node.

Open the Variable Definitions View to see these definitions:

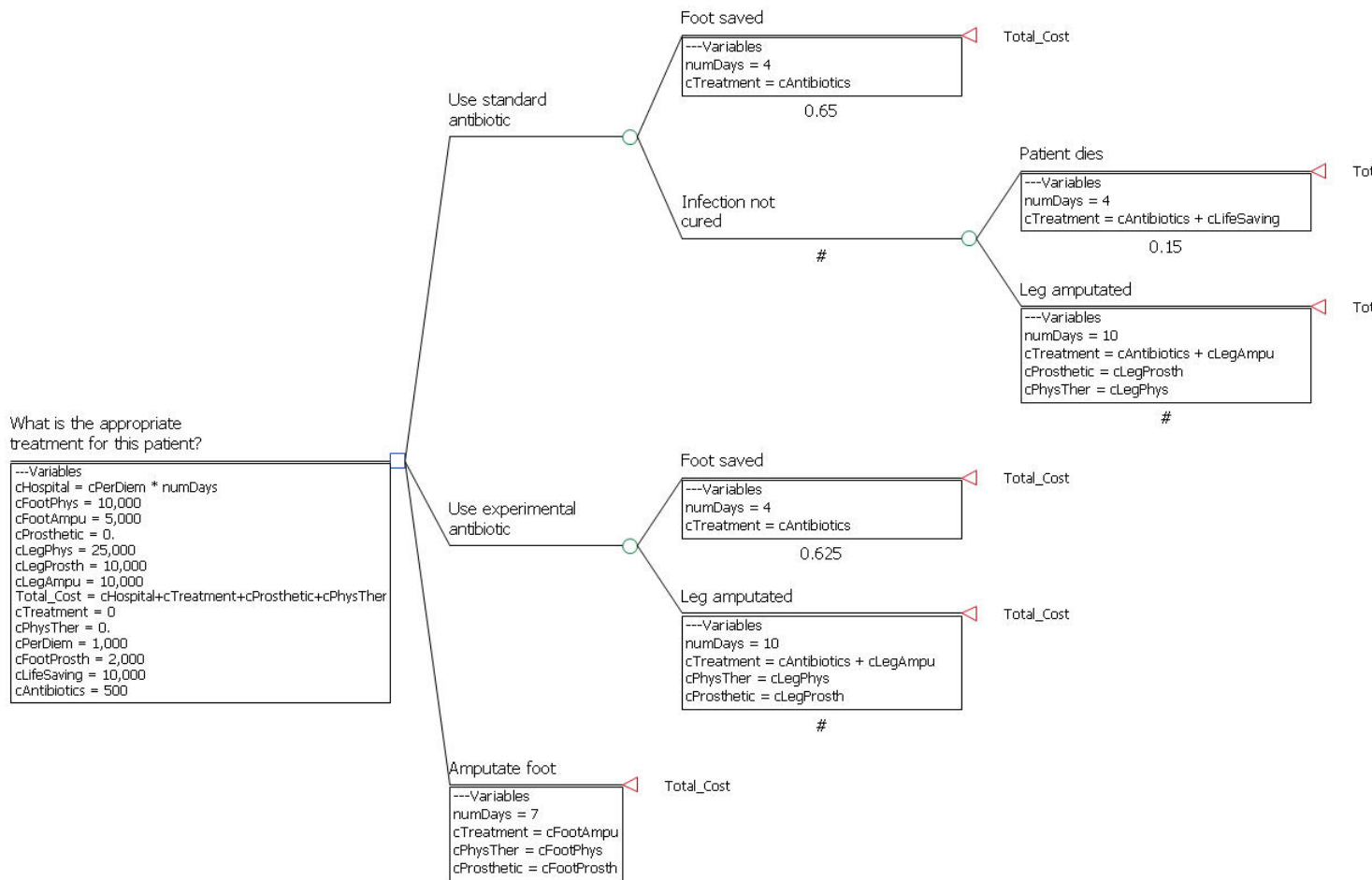
- Select the root node.
- Choose Node > Show View > Variable Definitions from the menu.

Much of the work required to make the tree more flexible has already been accomplished. All necessary variables have been created in the tree, and each has at least one definition.

The remaining tasks to be done in the tree are:

- assign Total_Cost as the payoff of every terminal node;
- redefine the variable (i.e., not fixed) components of Total_Cost for each outcome, as necessary.

The tree below illustrates how and where to add definitions of the variables used in the Total_Cost formula (other solutions are possible, as will be seen later in this chapter). Compare the tree you have opened to the picture of the completed tree.



Cost Formula tree with all variable definitions

17.7.3 Variables with multiple definitions

Note that, in the solution, several variables have multiple definitions. Note that, in the solution, several variables have *multiple definitions*. However, with the exception of numDays, no single variable has more than one numeric definition. As a general rule, it is advisable to avoid giving a single variable more than one numeric definition – for example, to represent the probability of two or more distinct events. It is important to follow this rule if you want to be able to perform sensitivity analysis on the variable in question.



The quantity represented by numDays, used in calculating cHospital, is not intended to be used in sensitivity analysis. If it were, separate variables would be used for each scenario.

17.7.4 Incorporate the formulas into the tree

Start by assigning the Total_Cost variable to all of the payoffs. Instead of deleting the existing numeric cost payoffs that are found in payoff #1, however, switch to payoff #2 and assign Total_Cost there.

To switch to payoff #2:

- Choose Tree > Tree Preferences from the menu or press *F11*.
- Navigate to the Tree Preferences category Calculation > Calculation Method > Simple.
- Change the Active payoff to 2.

All terminal nodes should now indicate that no payoff has been assigned. Now, update all the terminal nodes to use Total_Cost as the formula for payoff #2.

To change the payoffs:

- Double-click on the payoff expression next to the top terminal node.
- Enter "Total_Cost" as the expression for Payoff 2.
- Repeat for the remaining terminal nodes.

At this point, although the tree can calculate without causing errors, it will not calculate costs correctly because most of the components of Total_Cost are still defined equal to 0.

Examine the tree to see where and how variables are currently used and defined in the tree, and where definitions need to be added or modified.

To calculate the payoff of any node in the tree, TreeAge Pro must evaluate Total_Cost. To do this, it will search start searching for a definition of Total_Cost at the terminal node, and work leftward until it finds a definition. In this case, the first definition it will find is the default definition of Total_Cost from the root node, which accumulates the following components:

$(costofinpatientcare)+(costoftreatment)+(costofprosthetic)+(costofphysicaltherapy)$

The component variables cHospital, cTreatment, cProsthetic and cPhysTher need to be evaluated. TreeAge Pro restarts the search for each variable's definition *at the terminal node* which is being calculated. This right-to-left search is restarted for each variable encountered in the calculation. (The same process occurs for probability formulas, with a search rooted at the branch being calculated.)

Therefore, for every path in the tree, each variable in the cost formula must have an appropriate definition. Note that for some outcomes, some components do not apply. For example, there are three terminal nodes which represent non-amputation scenarios, and which therefore have no prosthetic or therapy costs. These components should have 0 values in these paths.

Incorporate the remaining variable definitions into the model. For example, at the *Amputate foot* node, you will need to add three additional variable definitions (for variables cTreatment, cPhysTher and cProsthetic).

You should confirm that, when rolled back, the Cost Variables tree provides the same results as the original tree. You can test this by changing the active payoff back and forth from 1 to 2.



There are two important exceptions to the right-to-left search rule. One involves the recursive definitions of variables, which will be covered at the end of this chapter. The other exception involves the evaluation of tracker variables; refer to the Healthcare Module documentation for details.

All payoff values are set to the variable `Total_Cost`, but the value are different for each terminal node. Let's examine one of the payoff calculations in detail, specifically at the *Amputate foot* node.

The Variable Definitions View for the node shows all the definitions we need to calculate the value of `Total_Cost`.

---Defined---	
cPhysTher	cFootPhys
cProsthetic	cFootProsth
cTreatment	cFootAmpu
numDays	7
---Undefined---	
---Inherited---	
What is the approj	
cAntibiotics	500
cFootAmpu	5,000
cFootPhys	10,000
cFootProsth	2,000
cHospital	cPerDiem * numDays
cLegAmpu	10,000
cLegPhys	25,000
cLegProsth	10,000
cLifeSaving	10,000
cPerDiem	1,000
cPhysTher	0.
cProsthetic	0.
cTreatment	0
Total_Cost	cHospital+cTreatment+cProsthetic+cPhysTher

Variable definitions at Amputate foot node

Based on the right-to-left lookup rule, `Total_Cost` is first defined at the root node.

$$Total_Cost = cHospital + cTreatment + cProsthetic + cPhysTher$$

In order to do that calculation, the individual components need to be calculated based on their closest definitions.

`cHospital` is calculated from the `cPerDiem` definition at the root node and the `numDays` definition at the current node.

$$cHospital = cPerDiem * numDays = 1,000 * 7 = 7,000$$

`cTreatment` is calculated from its definition at the current node, which in turn requires a definition of `cFootAmpu` from the root node.

$$cTreatment = cFootAmpu = 5,000$$

`cProsthetic` is calculated from its definition at the current node, which in turn requires a definition of `cFootProsth` from the root node.

$$cProsthetic = cFootProsth = 2,000$$

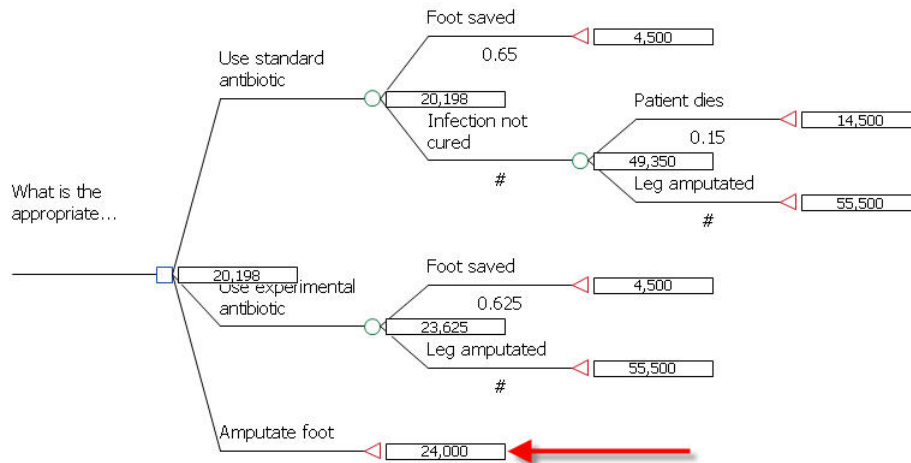
$cPhysTher$ is calculated from its definition at the current node, which in turn requires a definition of $cFootPhys$ from the root node.

$$cPhysTher = cFootPhys = 10,000$$

Now we can see the full calculation of the $Total_Cost$ value.

$$Total_Cost = 7,000 + 5,000 + 2,000 + 10,000 = 24,000$$

If you run roll back and check the payoff at the *Foot amputated* node, the value is \$24,000.



Cost Formula tree rolled back

17.8 Using mathematical, statistical, and other functions

TreeAge Pro includes a wide variety of functions which make it easier to perform some commonly-used mathematical operations, or in some cases to provide access to special features in TreeAge Pro.

- TreeAge Pro's built-in functions are case-insensitive, with the name followed by parentheses, and in most cases take arguments in the parentheses.
- Functions that have multiple arguments must use semicolon (",") separators.
- In most cases, any valid expression can be used as a function argument (with the exception of MatrixMult).
- In the tree window, use the Formula Editor for help in assigning the correct parameters to each functions.
- Functions which here indicate an argument named "LIST" take a flexible number of arguments. For example, Average() returns the arithmetic mean of all of its arguments, so Average(1;4;8;13) = 6.5.
- To test a function, use the Calculator/Evaluator tool described in Chapter 13.
- The Distributions Chapter covers a special set of functions used in Monte Carlo simulation.
- The Excel Linking Chapter covers a special set of functions that take text string arguments, instead of numeric or variable expressions.

- The Markov Modeling Tools Chapter covers a special set of functions used in Markov modeling, for example in calculating transition probabilities.

Notes on using functions in formulas

17.9 Arithmetic functions

Function	Explanation
Abs(x)	Absolute value of x
Average(LIST)	Arithmetic mean of a LIST of values/expressions
Ceiling(x)	Smallest integer larger than x
Exp(x)	“e” to the xth power e^x
Floor(x)	Greatest integer smaller than x
GammaFn(x)	(n-1)! for integers less than 19; Stirling's approximation otherwise. For factorial use GammaFn(x+1)
Int(x)	Integer component of x
Ln(x)	Natural (base “e”) logarithm of x
Log(x)	Base 10 logarithm of x
Max(LIST)	Maximum, or highest value, of a LIST
Min(LIST)	Minimum, or lowest value, of a LIST
Modulo(x; y)	Remainder of x divided by y
Prod(LIST)	Product of a LIST
Root(x; y)	yth root of x $\sqrt[y]{x}$
Round(x)	x rounded to the nearest integer
Sqrt(x)	Square root of x \sqrt{x}
Stdev(LIST)	Standard deviation of a LIST of numbers
Sum(LIST)	Sum of a list

Arithmetic functions



In the arithmetic functions, “e” represents the base of the natural logarithm (approximately 2.718.)

17.10 Financial/discounting functions

Function	Explanation
Annuity(rate; periods)	To calculate the net present value of a series of equal future payments, multiply this function times the amount of a single payment. $(1 - (1 + rate)^{-periods})/rate$
Compound(rate; periods)	Returns the compound interest rate (effective yield) at a fixed rate over a fixed number of periods. If used as a multiplier, this function can be used to calculate future value; if used as a divisor, it can be used to calculate discounted, present value. $(1 + rate)^{periods}$
Discount(util; rate; time)	Discounts a specified value (cost or utility) at the specified discount rate over the specified period. $util/((1 + rate)^{time})$
DiscountCont(util; rate; time_start; time_end) DiscountContDES(util; rate)	Discounts a specific value (cost or utility) at the specified rate on a continuous basis for DES models. DiscountCont requires that you provide the start and end time. DiscountContDES uses the defaults of _prior_time and _time for the start and end time. This only works correctly if the time measurement is annual.
FV(pmt; pv; rate; payments)	Returns the future value of a series of equal, periodic payments. The “pv” parameter represents an initial payment. $[pmt * ((1 + rate)^{payments} - 1)]/rate + pv * (1 + rate)^{payments}$
NPV(rate; LIST of flows)	Returns the net present value of periodic cash flows, discounted. $(f_i/(1 + rate)^i)$
PMT(principal; rate; term)	Returns the size of equal, periodic payments required to pay off a loan, given the principal, interest rate, and term of the loan. $principal * (rate/(1 - (1 + rate)^{-term}))$
PV(pmt; fv; rate; payments)	Returns the present value of equal, periodic payments at a fixed interest rate. The “fv” parameter represents a final payment. $[pmt * (1 - (1 + rate)^{-payments})]/rate + fv/((1 + rate)^{payments})$
UtilDiscount()	Obsolete. See Discount(), above.

Financial/discounting functions



In functions, a “rate” argument can be entered as either a percent or a decimal. For example, eight percent can be represented as either “8%” or “.08,” but not simply as “8.” Do not enter a very small percentage as “.08%”. However, it is OK to enter “0.08%”. References to periods/payments specifically mean the number of periods/number of payments.

17.11 Miscellaneous functions

Function	Explanation
Bilink(index)	Returns the value associated with a dynamic link. Refer to the Excel Linking Chapter.
BranchProb()	Returns the calculated branch probability for the node being calculated. Unlike PathProb(), this function only includes the probability at the specific branch node.
Choose(index;LIST of values)	Returns a value based on its location in the LIST, as specified by the index. The index must be a positive integer; an error is reported for fractional or out-of-range values. For example, “ <i>Choose(2;100;200;300)</i> ” returns 200, because the index is 2, and 200 is the second value in the list. This function can often be used to replace complex nested <i>If</i> functions.
If(condition; trueval; falseval)	Evaluates a condition and returns “trueval” if the condition is true or “falseval” if the condition is false. For example, “ <i>If(x<0;50;75)</i> ” would return 50 if x were negative and 75 if x were non-negative.
Inf()	Returns infinity (#). Although no arguments are accepted, you must type the left and right parentheses.
Link(index)	Returns the value associated with a DDE link. See Bilink() above.
Pi()	Returns #, or approximately 3.1416.
PathProb()	Returns the cumulative path probability for the node being calculated. In a Markov subtree, PathProb() returns the cumulative path probability from the Markov state to the node being calculated. In a non-Markov tree, you can use this function to calculate the percentage of an overall cohort that reaches a specific node.
StateProb(A; B)	Returns the state probability of one or more states at the start of the current cycle. Refer to the Markov Modeling Tools Chapter for details on this Markov modeling function.
Sub(index)	Obsolete. See Bilink() and Link(), above.

Miscellaneous functions



Choose() function tip: Performing a sensitivity analysis on a variable used as the index of the *Choose()* function, from 1 to the number of list items, can show the impact on calculations of the different values specified in the list.

Nesting functions: All functions can be nested. The *If()* function is frequently used nested within another *If()* function call. This is because the *If()* function tests for one condition, and can return either of two values, normally. However, by using another *If()* function as one of the return values, it is possible to perform two tests and return any of three values (and so on).



String functions: The Tools and Functions for Complex Trees Chapter covers a special set of functions that take text string arguments, instead of numeric or variable expressions.

Markov functions: The Markov Modeling Tools Chapter covers a special set of functions used in Markov modeling, for example in calculating transition probabilities.

17.12 Using mathematical, statistical, and other functions

- TreeAge Pro's built-in functions are case-insensitive, with the name followed by parentheses, and in most cases take arguments in the parentheses.
- Functions that have multiple arguments must use semicolon (";") separators.
- In most cases, any valid expression can be used as an function argument (with the exception of *MatrixMult*).
- A separate section at the end of the chapter covers the special set of functions that take text string arguments, instead of numeric or variable expressions.
- Functions which here indicate an argument named "LIST" take a flexible number of arguments. For example, *Average()* returns the arithmetic mean of all of its arguments, so *Average(1;4;8;13) = 6.5*.
- In the tree window, use the Function Helper (see Chapter 13) for help in assigning the correct parameters to each functions. The functions are described

17.13 Recursive variable definitions

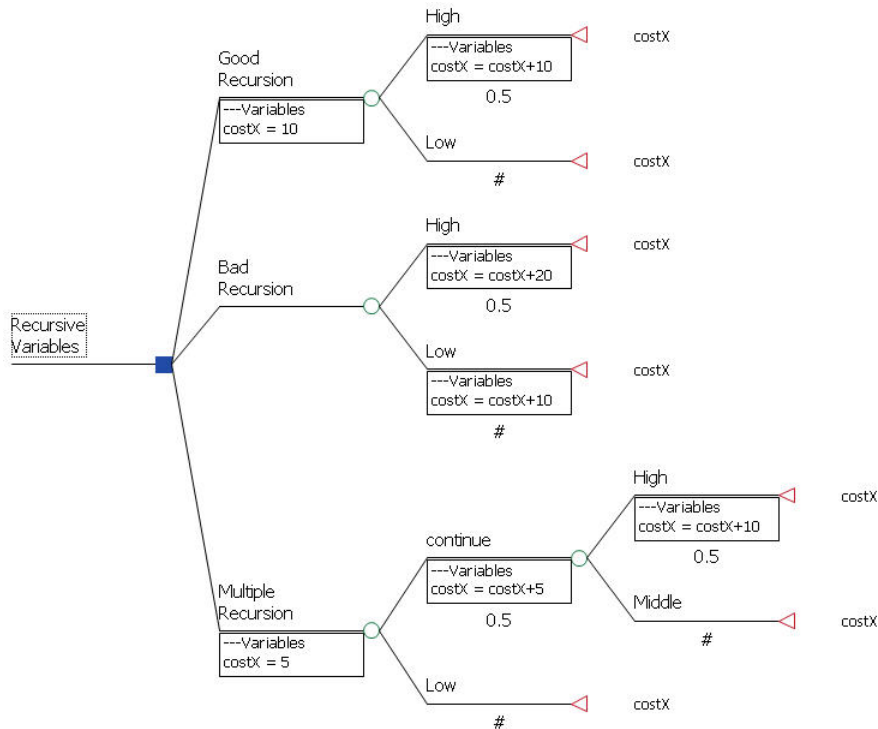
A recursive variable definition means a definition in which the variable being defined also occurs in the definition. Recursive definitions can be an effective way to build complex cost formulas, like those in the Cost Variables tree. Rather than creating one or more long formulas, you can gradually build the formula, adding components as events occur in each particular scenario.

17.13.1 How recursive definitions work

During calculation of a probability, payoff, or Markov reward, when the standard right-to-left search for a definition of a particular variable (e.g., "costX") first locates a definition (e.g., "costX=1,000"), TreeAge Pro stops looking for additional definitions of that variable. However, when the first definition TreeAge

Pro encounters in the search is a self-referential, recursive definition (e.g., “costX=costX+1,000”), the variable (e.g., “costX”) is flagged as a recursive variable, and TreeAge Pro continues searching for additional definitions of that variable to the left of the node where the first, recursive definition was found.

Open the Special Features tutorial example tree Recursive Variables, shown below, to see an example of this process.



Recursive Variables model

All payoffs in the tree reference the same variable, costX, which has no default definition. The decision node's topmost subtree, labeled Good Recursion, illustrates a valid recursive definition. When calculating the payoff of the first terminal node in the Good Recursion subtree, labeled High, the normal, right-to-left search for a definition of costX finds the self-referential definition costX=costX+10 at that terminal node.

For the purposes of the current terminal node payoff calculation, costX is now identified as a recursively defined variable. The search for additional definitions of costX is now continued one node to the left, at the Good Recursion node. There, the non-recursive definition costX=10 is found, the search is complete, and the payoff calculation can be carried out.

Select the terminal node labeled High in the Good Recursion subtree, and choose Analysis > Expected Value. The calculated value is 20.

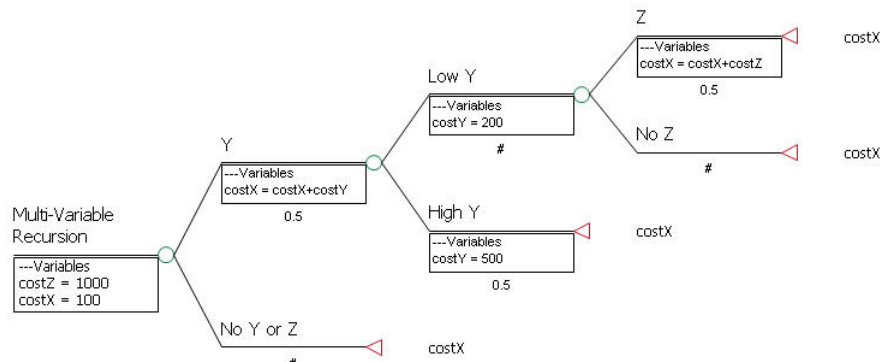
A non-recursive definition of the recursive variable, like costX=10 at the Good Recursion node, must eventually be found; definitions can't be infinitely recursive.

Look at the second branch of the decision node, labeled Bad Recursion. In this subtree, when either terminal node's payoff is evaluated, TreeAge Pro's variable definition search locates a recursive definition. In both cases, the search for additional definitions is started one node to the left, at the Bad Recursion node. No non-recursive definition of x has been made there or at the root node. Therefore, if you try to calculate the Bad Recursion subtree, an error message will be shown. To see this, select the Bad Recursion node or either of its terminal nodes and choose Analysis > Expected Value command.

Multiple recursions, using a series of recursive definitions of a variable, will work, as shown in the Multiple Recursion subtree. Simply ensure that a numeric definition of the recursive variable will eventually be found. For example, in calculating the payoff of the terminal node labeled Middle in the Multiple Recursion subtree, the variable definition search locates the recursive definitions $\text{costX} = \text{costX} + 10$ and $\text{costX} = \text{costX} + 5$, and finally the non-recursive definition $\text{costX} = 0$. Thus, the calculated value of the Middle terminal node is 15.

17.13.2 Complex recursion

Other variables may be referenced in a recursive definition. To see how TreeAge Pro's variable definition search works with a combination of recursive and non-recursive variables, open the tutorial example tree "Multi-Variable Recursion".



Multi-Variable Recursion

To better explain this complex example, the text will illustrate the incremental changes in the payoff calculation formula during TreeAge Pro's variable definition search.

All payoffs in the Multi Variable Recursion tree use the same variable, costX (payoff = costX). When the terminal node labeled Z is evaluated, the variable definition search finds the recursive definition $\text{costX} = \text{costX} + \text{costZ}$ at the terminal node (payoff = $\text{costX} + \text{costZ}$).

Before a recursive search for costX is continued one node to the left, TreeAge Pro looks for a definition for costZ . The variable costZ is evaluated as a normal variable, which means that the search for a definition of costZ is started at the node being calculated, the terminal node Z. A normal right-to-left search locates the non-recursive definition $\text{costZ} = 1000$ at the root node (payoff = $\text{costX} + 1000$).

The recursive search for additional definitions of costX is then continued at the Low Y chance node, which is one node to the left of the node where the initial, recursive definition of costX was found. Another recursive definition, $\text{costX} = \text{costX} + \text{costY}$, is found at the chance node labeled Y (payoff = $[\text{costX} + 1000] + \text{costY}$).

As above, before the recursive search continues, a normal variable definition search for costY is initiated at the terminal node being calculated, Z. At the chance node Low Y, the non-recursive definition $\text{costY} = 200$ is located (payoff = $[\text{costX} + 1000] + 200$).

TreeAge Pro proceeds with the search for additional definitions of costX, starting at the root node. A non-recursive definition of costX is required, and found: $\text{costX} = 100$. The final payoff formula for node Z, therefore, is $[100 + 1000] + 200$. Calculating an expected value for the Z terminal node returns the value 1300.

Before using complex recursion in your models, it is important that you be thoroughly familiar with the logic underlying both standard and recursive variable definitions in TreeAge Pro. You are also urged to test your model to make sure it appears to be calculating correctly.

In summary, the payoff value is costX ...

$$\text{payoff} = \text{costX}$$

Based on the recursive definition at node Z, the expression changes to...

$$\text{payoff} = \text{costX} + \text{costZ}$$

Substituting in the recursive definition for costX at node Y, the expression changes to...

$$\text{payoff} = (\text{costX} + \text{costY}) + \text{costZ}$$

Now the right-to left definitions for each variable can be resolved...

$$\text{payoff} = 100 + 200 + 1,000 = 1,300$$

The expected value at node Z is, in fact, 1,300.

17.14 Operators

Mathematical expressions used in TreeAge Pro necessarily contain operators (such as addition or multiplication signs). This section describes operators available in creating expressions.

17.14.1 Arithmetic operators

These operators perform arithmetic on the values that surround them. TreeAge Pro uses the traditional syntax for expressions, known as infix notation. For example, an expression that adds three and seven would be written $3 + 7$, rather than $3, 7 +$.

Symbol	Example	Description
+	$x + y$	Addition. Returns the sum of x and y.
-	$x - y$	Subtraction. Returns the difference between x and y. (Also used for negation, i.e., to denote negative numbers.)
*	$x * y$	Multiplication. Returns the product of x and y.
/	x / y	Division. Returns the quotient of x and y.
^	$x ^ y$	Exponentiation. Returns x to the yth power.
()	$x * (y+z)$	Grouping. Returns the product of x and the sum of y and z.

Arithmetic operators**17.14.2 Relational operators**

These operators return a true or false value, depending on the veracity of the expression in which they appear. A true value is represented by a numeric 1, a false value receives a numeric value of 0. Relational operators are useful in many settings: in If() and Choose() functions, in expressions evaluated at logic nodes, and in a Markov termination condition.

Symbol	Example	Description
<	$x < y$	Less than. Returns true if x is less than y, and false if x is greater than or equal to y.
<=	$x <= y$	Less than or equal to. Returns true if x is less than or equal to y, and false if x is greater than y.
>	$x > y$	Greater than. Returns true if x is greater than y, and false if x is less than or equal to y.
>=	$x >= y$	Greater than or equal to. Returns true if x is greater than or equal to y, and false if x is less than y.
=	$x = y$	Equals. Returns true if x equals y, and false if x is not equal to y.
<>	$x <> y$	Not equal to. Returns true if x is not equal to y, and false if x equals y.

Relational operators

It is also possible, using the appropriate relational expression syntax, to test one value in terms of two others. There are a number of acceptable forms, with the two basic ones being:

- $y < x < z$ - Returns true if x is both (a) greater than y and (b) less than z.
- $y > x > z$ - Returns true if x is both (a) less than y and (b) greater than z.

Other valid forms of this syntax can be created by substituting ">=" for ">" or "<=" for "<" (for example, expressions of the form " $y <= x < z$ " and " $y >= x >= z$ " are valid). These are the only valid substitutions, though (for example, expressions of the form " $y < x > z$ " are not valid). Failure to follow

these rules when creating relational expressions of this kind will likely result in unintended calculation results.

17.14.3 Logical operators

Three logical operators are also available: logical AND, logical (inclusive) OR, and logical NOT.

AND is represented by the ampersand (&), OR by the vertical bar (|), and NOT by the exclamation mark (!). Like the relational operators (which return 1 if a comparison is true and 0 if not), these logical operators are zero-centric. That is, any operand that is non-zero is treated as true, and only a zero operand is treated as false. The returned value is 1 if the evaluation is true, and 0 if false.

Symbol	Example	Description
&	$w < x \ \& \ y < z$	Logical AND. Returns true if w is less than x AND y is less than z.
	$w < x \ \ y < z$	Logical OR. Returns true if either w is less than x OR y is less than z.
!	$!(w < x)$	Logical NOT. Returns true if w is not less than x.

Logical operators

17.14.4 Operator precedence

In most situations, you will not need to know the details of which operators bind most tightly. However, when formulas do not appear to calculate correctly, you should check this section to see if precedence is a factor.

Operator precedence is how TreeAge Pro decides where you intended to put parentheses. Consider the following example:

$$A + B * C + D$$

A quick check of the precedence list below indicates that multiplication has higher precedence (binds more tightly) than addition. TreeAge Pro will therefore interpret your expression as:

$$A + (B * C) + D$$

This process is continued until all uncertain bindings are resolved.

The table below lists the operators available in TreeAge Pro in order of precedence. Operators with higher precedence will bind more tightly. Adjacent operators having the same precedence value will be applied from left to right.

Operator	Character	Precedence Value
Unary minus	-	8
Logical NOT	!	8
Exponent	^	7

Operator	Character	Precedence Value
Multiplication	*	6
Division	/	6
Addition	+	5
Subtraction	-	5
Comparators	<, <=, etc.	4
Logical AND	&	3
Logical OR		2
Parentheses, Brackets	(), []	1
Functional argument separator	;	1

Operator precedence

Notice that parentheses are at the bottom of the list. This simply means that the operators inside the parentheses will bind tightly to stay within the parentheses. They are your most useful tool for indicating your particular precedence requirements.

17.15 Keywords

Keywords are reserved values auto-generated by TreeAge Pro that can be accessed within your model. These keywords are used only in Markov models.

Keyword	Description
_stage	Markov cycle counter starting at 0
_tunnel	Markov cycle-in-state counter starting at 1
_trial	Trial iteration within Microsimulation
_sample	Sample iteration within PSA simulation
_trial_size	Total number of iterations for Microsimulation
_sample_size	Total number of iterations for PSA simulation
_voi_sample	Sample iteration for outer loop of EVPPI simulation
_voi_sample_size	Total number of iterations for outer loop of EVPPI simulation
_parallel_trials_creator	New trial iteration generated in parallel trials Microsimulation
_node_id	Return the numeric identifier for a specific node
_node_label	Return the label for a specific node

TreeAge Pro Keywords

18. More Sensitivity Analysis Tools

The last 3 chapters provided detailed instructions on representing uncertain values using variables and performing one-way sensitivity analysis. This chapter covers sensitivity analysis options, multi-way sensitivity analysis, and special variations on one-way sensitivity analysis.

18.1 Analyzing variables with multiple definitions

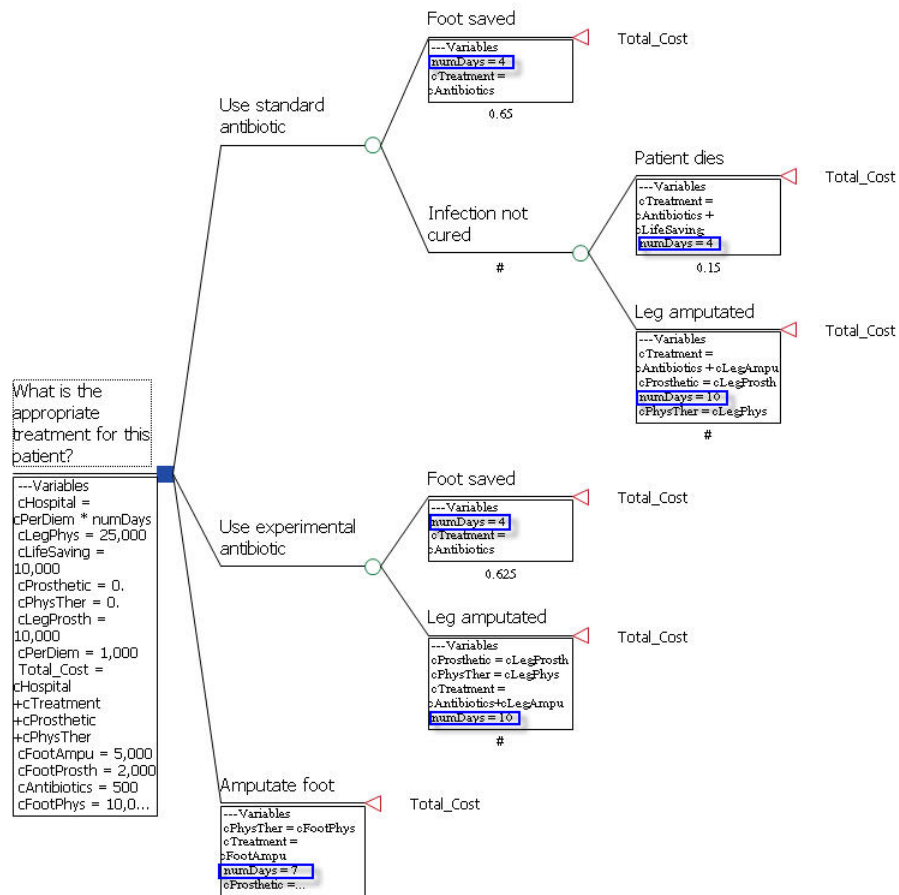
While first learning how to use variables, it is not uncommon to unintentionally end up with a variable defined numerically at multiple nodes. While there are good reasons why TreeAge Pro allows the same variable to be defined at multiple nodes — as described in prior chapters — this situation is usually neither necessary or desirable with a variable intended for sensitivity analysis, as it may result in errors.



It is strongly recommended that sensitivity analysis be limited to parameters defined at the root node with a numeric value. Variables that are defined by other variables, functions, etc. or that are defined at multiple nodes are generally not good candidates for sensitivity analysis. It is almost always possible to restructure variable definitions in such a way that the variable in question can be isolated as a numeric value at the root node.

We recommend that you avoid running sensitivity analysis on a variable with multiple definitions. However, this technique is shown for *illustration only*. If you must run sensitivity analysis on a variable with multiple definitions, you must specify which definitions should be used/changed within the sensitivity analysis.

We will look at running sensitivity analysis on the Get Started tutorial example tree, Climber Cost, specifically on the variable *numDays*.



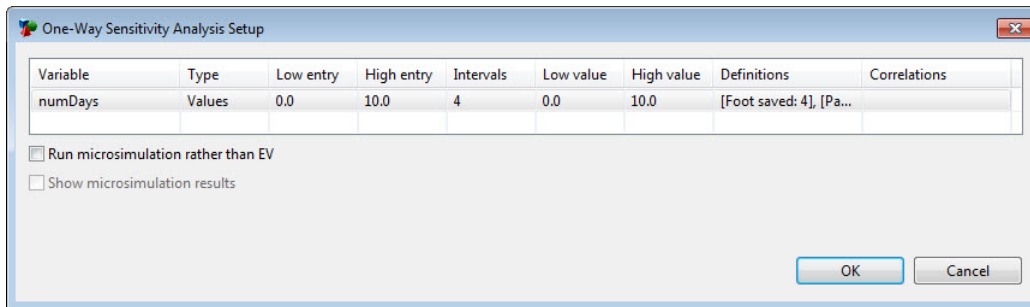
Climber Cost tree

Note that there are several definitions for the variable numDays. You might want to run sensitivity analysis specifically on the number of days in the hospital associated with leg amputation. In such a case, you would only want to use the numDays variable definitions at the two nodes labeled Leg amputated.



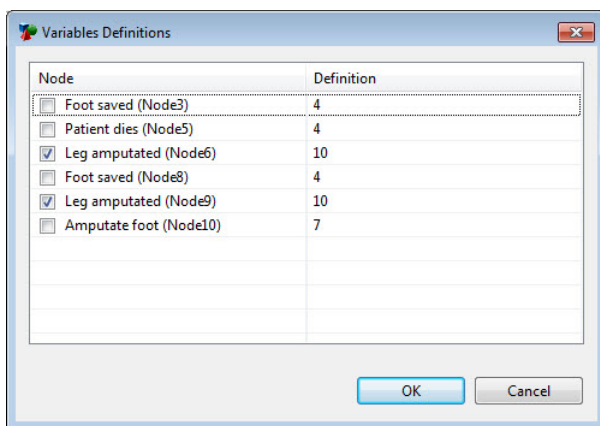
A better and safer way to achieve this result would be to define separate numeric parameters numDaysLegAmpu, numDaysFootAmpu, etc. at the root node. Then you could set the generic numDays variable equal to the appropriate numeric parameter at different nodes in the tree. Then you could run sensitivity analysis on the specific numeric parameter in question.

After selecting the variable numDays for sensitivity analysis in the One-Way Sensitivity Analysis Setup Dialog, click on the elipsis in the Definitions column.



Sensitivity analysis setup dialog

You are then presented with another dialog to select the specific nodes where you want the variable definitions used for sensitivity analysis.



Sensitivity analysis setup - select nodes

Note that only the two Leg amputated nodes are selected. When sensitivity analysis runs, only the definitions at those two nodes will be altered.

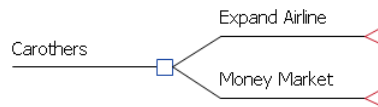
18.2 Tornado diagrams

A tornado diagram is a set of one-way sensitivity analyses brought together in a single graph. It can include any number of the variables defined in the tree.

In the graph, a horizontal bar is generated for each variable being analyzed. Expected (or incremental) value is displayed on the horizontal axis, so each bar represents the selected node's range of expected (or incremental) values generated by varying the related variable. A wide bar indicates that the associated variable has a large potential effect on the expected value of your model (given the range provided for each variable).

The graph is called a tornado diagram because the bars are arranged in order, with the widest bar (potentially the most critical uncertainty) at the top and the narrowest bar at the bottom, resulting in a funnel-like appearance.

The Other tutorial example tree, Airline Problem, is ready for a tornado diagram. The model is a simple cost function, each of whose inputs may be varied to see how each may affect the expected value.



Airline Problem tree

To create a tornado diagram:

- Select the decision node.
- Choose Analysis > Sensitivity Analysis > Tornado Diagram.
- In the Tornado Diagram setup dialog, select each variable to analyze and enter the variable's low and high values and intervals (see below).
- Click OK to start the analysis.

Variable	Type	Low entry	High entry	Intervals	Low value	High value	Definitions	Correlations
Cap_Sched	Values	0.4	0.6	4	0.4	0.6	[Carothers: .5]	
Charter_Price	Values	300.0	350.0	4	300.0	350.0	[Carothers: 325]	
Charter_Ratio	Values	0.45	0.7	4	0.45	0.7	[Carothers: 0.5]	
Hours_Flown	Values	500.0	1,000.0	4	500.0	1,000.0	[Carothers: 800]	
Insurance	Values	18,000.0	25,000.0	4	18,000.0	25,000.0	[Carothers: 20000]	
Interest_Rate	Values	0.105	0.13	4	0.105	0.13	[Carothers: .115]	
PctFinanced	Values	0.3	0.5	4	0.3	0.5	[Carothers: .4]	
Price	Values	85,000.0	90,000.0	4	85,000.0	90,000.0	[Carothers: 87500]	
Ticket_Price	Values	95.0	108.0	4	95.0	108.0	[Carothers: 100]	

☒ Extend bars using threshold info

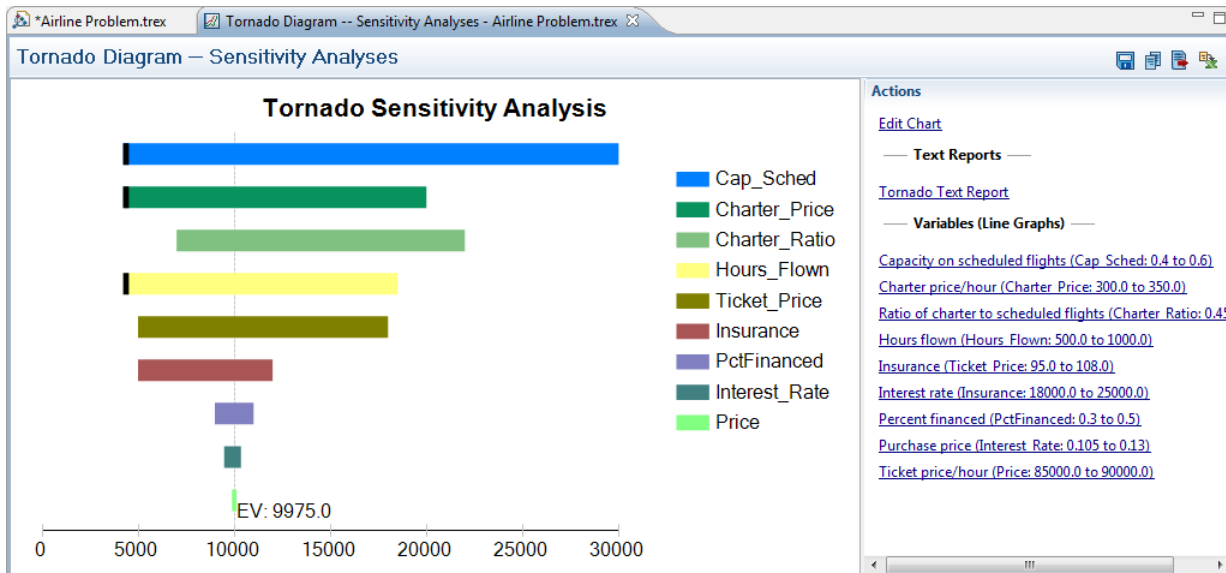
OK Cancel

Tornado Diagram setup



Be careful not to select variables that are formulas (i.e., Total_Cost, Total_Revenue), rather than numeric parameters.

The tornado diagram for the airline decision is shown below.



Tornado Diagram

Each bar represents a one-way sensitivity analysis performed at the selected node. The tornado diagram includes a vertical dotted line indicating the expected value. You can use this as a visual fulcrum, to view the impact of each variable relative to the original (baseline) expected value.

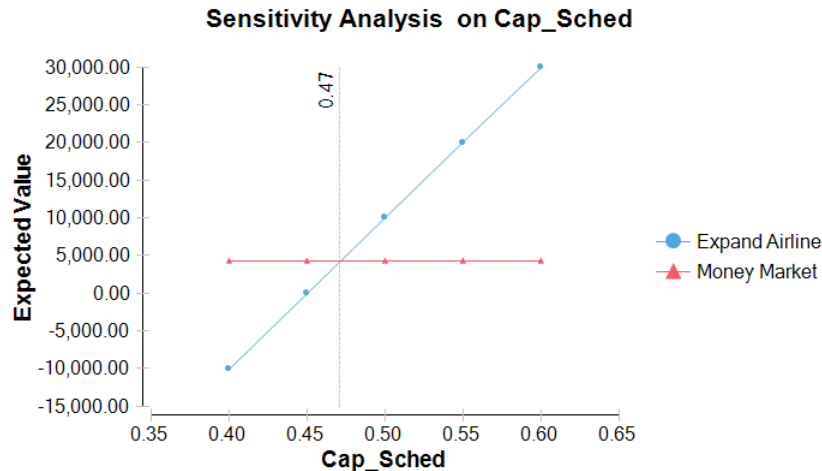
If you click on the Tornado Text Report link to the right of the graph, you will see the input and output range for each parameter.

VARIABLENAME	VARIABLERANGE	LOWVALUE	HIGHVALUE	SPREAD	SPREADSQR	RISKPCT	CUMULPCT
Cap_Sched	0.4 to 0.6	4,200	29,975	25,775	664,350,625	0.353	0.353
Operating_Cost	230.0 to 260.0	4,200	21,975	17,775	315,950,625	0.168	0.521
Charter_Price	300.0 to 350.0	4,200	19,975	15,775	248,850,625	0.132	0.654
Charter_Ratio	0.45 to 0.7	6,975	21,975	15,000	225,000,000	0.12	0.773
Hours_Flow	500.0 to 1000.0	4,200	18,475	14,275	203,775,625	0.108	0.882
Ticket_Price	95.0 to 108.0	4,975	17,975	13,000	169,000,000	0.09	0.971
Insurance	18000.0 to 25000.0	4,975	11,975	7,000	49,000,000	0.026	0.997
PctFinanced	0.3 to 0.5	8,968.75	10,981.25	2,012.5	4,050,156.25	0.002	1
Interest_Rate	0.105 to 0.13	9,450	10,325	875	765,625	0	1
Price	85000.0 to 90000.0	9,860	10,090	230	52,900	0	1

Tornado Diagram Text Report

The variable range column shows the range you entered for each variable. The low and high values show the the lowest and highest expected value for the model based on the *optimal strategy at each point*. Note that none of the values in the low value can reach below \$4,200, since that is the fixed value for the *Money Market* strategy. The remaining columns are described later in this section.

If you expand the Variables group to the right of the graph, you are presented with a link for each variable included in the Tornado diagram. Click on any of those links to see the One-Way Sensitivity Analysis graph associated with that variable. See the *Cap_Sched* graph below.



One-Way Sensitivity Analysis Graph - from tornado diagram link

Tornado diagrams can be created at chance nodes and decision nodes. At a decision node, any threshold (i.e., change in policy) found will be identified in a variable's tornado bar with a heavy vertical line. Threshold lines are drawn at the expected value on the x-axis at which the optimal path changes. Note the heavy vertical line at the left end of the *Cap_Sched* bar of the Tornado Diagram. This indicates a change in strategy to the *Money Market* strategy (also shown in the one-way graph above). Since that strategy is fixed at \$4,200, none of the bars can extend lower than that.

If a threshold appears at either end of a bar, this usually indicates that an alternative which is optimal for part of the analysis range has an unchanging expected value in that range. To see details about the change in policy associated with a threshold line, open the One-Way Sensitivity Analysis graph associated with that variable.



Because setting up the tornado diagram analysis can be time consuming, you may want to store the setup information. Refer to the Stored Analysis Abstracts and Sequences Chapter for details on using stored analyses.

18.2.1 Incremental calculations

A tornado diagram can report, instead, the sensitivity of the incremental/marginal value calculated between two strategies.

To create an incremental tornado diagram:

- Run a regular tornado diagram as described above.
- Expand the Incremental Tornado... group label to the right of the tornado diagram.
- Click on the link representing the two strategies you want to compare.

Note that every combination of strategies will be listed as an option, including two links for each pair switching the comparator and baseline.

18.2.2 Additional calculations in the text report

The tornado diagram's text report will display, in addition to the input and output ranges for each parameter, a number of other useful calculated values.

- *Spread* – This is the width of the bar (i.e., High EV - Low EV).
- *SpreadSqr* – The spread value, squared.

Adding the SpreadSqr values to calculate a net risk value, two additional measures of uncertainty are then calculated for each variable.

- *Risk Pct* – This is a measure of how much of the total uncertainty is represented by the specified bar (equals SpreadSqr / NetRisk). The RiskPct values sum to 1.0.
- *Cum Pct* – A cumulative version of Risk Pct., making it easy to scan the bars and say “to address 90% of the risk, I must consider the uncertainty represented by the following variables...”

18.2.3 Including correlated variables in the tornado diagram

As in all sensitivity analyses, if you select a variable for the tornado diagram which has correlations, the correlated variables will vary together. The Analyzing correlated variables section of this chapter contains additional information related to sensitivity analysis on correlations.

The names of the correlated variables are shown in the list of variables available for inclusion in a Tornado Diagram. A pair of correlated variables cannot both be selected for the Tornado Diagram as they would have identical results.

18.3 Two-way sensitivity analysis

Two-way sensitivity analysis is used to examine the impact on a decision of simultaneous changes in the values of two variables. One method is to run a series of one-way analyses, each time incrementing the value of a second variable.

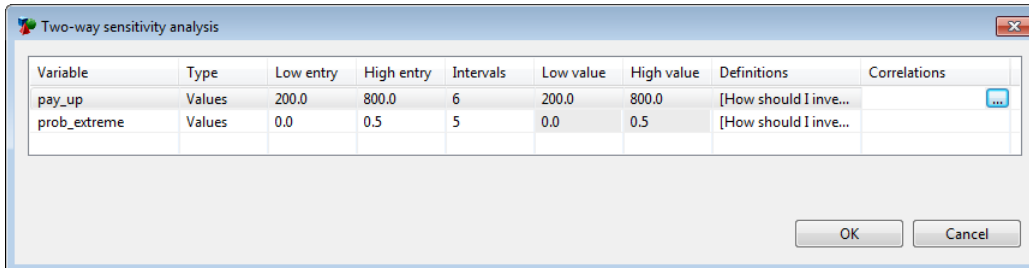
Another method available in TreeAge Pro is to automate this series of analyses and present the results in a region graph. The region graph very efficiently identifies changes in the optimal policy as the values of the two variables change.

The two-way sensitivity analysis setup dialog resembles the one-way dialog, except that you must specify two variables and a range of values for each. A decision node must be selected in order to perform a two-way sensitivity analysis.

To perform a two-way sensitivity analysis:

- Open the Get Started tutorial example tree, Three Variables.
- Select the root node.

- Choose Analysis > Sensitivity Analysis > Two-Way.
- For one variable, select pay_up, and specify the range from 200 to 800 with 6 intervals.
- For the second variable, select prob_extreme, and specify the range from 0 to 0.5 with 5 intervals
- Click OK to run the analysis.



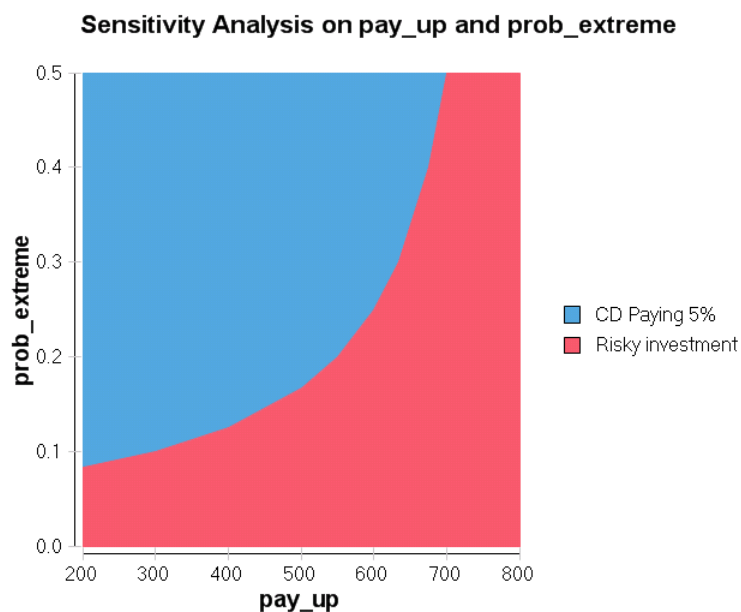
The dialog box titled "Two-way sensitivity analysis" contains a table with the following data:

Variable	Type	Low entry	High entry	Intervals	Low value	High value	Definitions	Correlations
pay_up	Values	200.0	800.0	6	200.0	800.0	[How should I inve...	...
prob_extreme	Values	0.0	0.5	5	0.0	0.5	[How should I inve...	

At the bottom right are "OK" and "Cancel" buttons.

Two-Way Sensitivity Analysis Setup

The resulting graph is shown below. It identifies which strategy is optimal in regions of values of the variables; thresholds are simply the border between two regions.



Two-Way Sensitivity Analysis Graph

The first variable in the setup dialog (pay_up) forms the horizontal axis of the graph, while the second variable (prob_extreme) forms the vertical axis.



It is recommended that you use two-way analysis *only* when the two variables are independent. If the two variables are a correlated pair, be sure to turn off the correlation.

18.3.1 Choosing intervals

Based on the selecting X intervals for one variable, and Y intervals for the other, TreeAge Pro recalculates the tree $(X+1)*(Y+1)$ times, for the different combinations of values of the two variables. In the example on the previous page, 42 (7x6) recalculations occur, creating a grid of sensitivity analysis results.

You may find that a two-way analysis requires more intervals per variable to attain a reasonable level of accuracy than a one-way analysis. This is because a two-way analysis graphically represents only the threshold values – the optimal path crossings. The one-way analysis may show significant details which are simply not shown in the two-way analysis.

The graphical representation of the results of two-way sensitivity analysis has some unavoidable limitations; this also applies to three-way analysis.

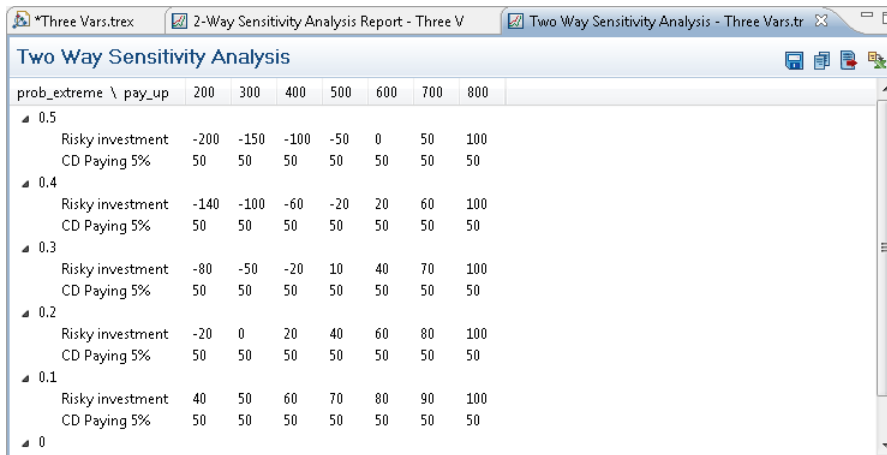


The accuracy of threshold lines may be compromised around the edges of the graph. The unavoidable result of using approximation techniques to identify thresholds is the appearance of distortion when two edges of a region of optimality draw closer together than one-half the width of an axis interval. Accuracy can be enhanced by running the analysis using more intervals.

Threshold lines represent points of indifference. Regions of indifference, however, are not shown. Areas of the graph where indifference exists are instead assigned to one alternative. You should use the text report (accessed via the Actions > or Graph > Text Report... command) to identify any areas of indifference by comparing the expected values at each interval. Isocontours (next section, but not yet implemented.) of small magnitude may also help identify where indifference ends.

18.3.2 Two-way sensitivity analysis text report

The region graph's text report is accessible from the "Text Report" link to the right of the graph. The text report displays a grouped matrix of results.



Two-Way Sensitivity Analysis Text Report

The text report's layout is based on the graph axes. The variable from the vertical graph axis is represented as grouped rows in the report; the variable from the horizontal graph axis is represented by the columns of the report. For each combination of values from the two variables, a list of expected values is reported for the branches of the decision node.

To change the orientation of the two variables, rerun the analysis with the other variable selected first.

18.3.3 Custom isocontours

Isocontours have not yet been implemented in TreeAge Pro 201x.

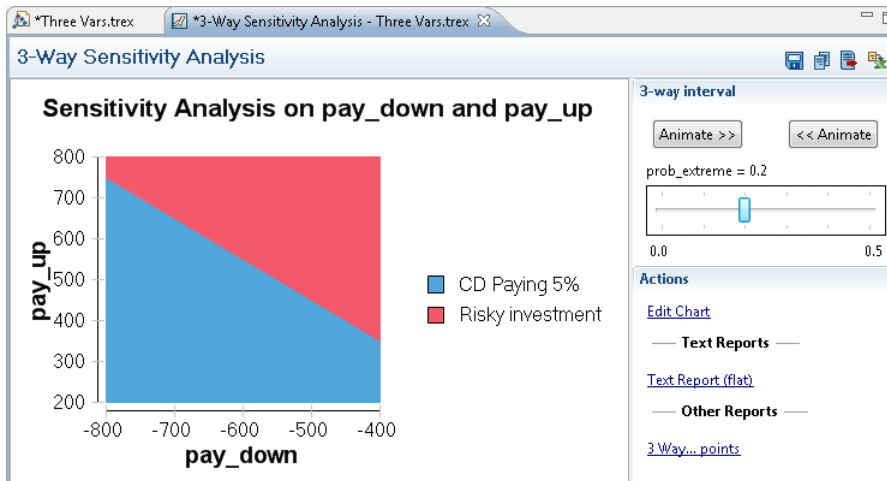
18.4 Three-way sensitivity analysis

The three-way sensitivity analysis dialog looks the same as the two- and one-way sensitivity analysis dialog except that three variable ranges are entered.

Three-Way Sensitivity Analysis Setup

Three variables cannot be presented clearly in a two-dimensional graph. Therefore, the results of a three-way sensitivity analysis are presented as an animated two-way sensitivity analysis region graph.

The third variable is represented not with its own axis, but rather using a series of two-way graphs — if four intervals are specified for the third variable, then five graphs will be created, and shown in series.



Three-way sensitivity analysis

Use the Animate button or the scroll bar to cause the third variable to cycle through its range. The successive frames of the three-way analysis shows how the two-way region/optimality graph for the first two variables is affected by varying the value of the third variable.

18.4.1 Three-way sensitivity text report

The three-way version of the region graph now allows viewing and exporting the text report for all combinations of variables. Since three variable inputs and the corresponding outputs cannot easily be presented in a grid, only the "flat" version of the Text Report is available.

VARIABLE1	VARIABLE2	VARIABLE3	STRATEGYINDEX	STRATEGY	VALUE
-400	800	0.5	0	Risky investment	200
-400	800	0.5	1	CD Paying 5%	50
-400	800	0.4	0	Risky investment	180
-400	800	0.4	1	CD Paying 5%	50
-400	800	0.3	0	Risky investment	160
-400	800	0.3	1	CD Paying 5%	50
-400	800	0.2	0	Risky investment	140
-400	800	0.2	1	CD Paying 5%	50
-400	800	0.1	0	Risky investment	120
-400	800	0.1	1	CD Paying 5%	50
-400	800	0	0	Risky investment	100
-400	800	0	1	CD Paying 5%	50
-400	700	0.5	0	Risky investment	150
-400	700	0.5	1	CD Paying 5%	50
-400	700	0.4	0	Risky investment	140
-400	700	0.4	1	CD Paying 5%	50

Three-way sensitivity analysis text report

The report includes the expected value for each strategy for every combination of the three variables.

18.5 Threshold analysis

This specialized form of sensitivity analysis offers the ability to search more thoroughly and accurately for threshold information. The result of this analysis is a detailed, textual description of how the optimal strategy is affected by changing the value of a single variable across a designated range.

In a standard one-way sensitivity analysis, the user designates the number of intervals into which the range is to be divided; actual calculations occur only at these intervals. As a result, the accuracy of the associated threshold analysis is limited to values determined by linear interpolation.

In contrast, threshold analysis has been designed to maximize accuracy of the analysis in situations where accuracy is more critical than speed. The specified range is iteratively searched until a specified minimum tolerance is reached.

This section will use the Get Started tutorial example tree Three Variables to illustrate threshold analysis.

To run threshold analysis:

- Choose Analysis > Sensitivity Analysis > Threshold Analysis from the menu.
- In the Threshold Analysis dialog, specify a variable and range, a tolerance and a degree of linearity. See below.

Variable	Type	Low en...	High e...	Intervals	Low val...	High v...	Definitions	Correlations
prob_extreme	Values	0.0	0.5	4	0.0	0.5	[How should I ...]	

☐ Run microsimulation rather than EV

☐ Show microsimulation results

Willingness-to-pay:

Tolerance:

Degree of non-linearity

☒ Low (4)

☐ Medium (8)

☐ High/don't know (12)

Tolerance uses same units as variable range.
Non-linearity setting controls number of sub-intervals used to reanalyze intervals.

OK Cancel

Threshold Analysis Dialog

The variable selection is the same as for one-way sensitivity analysis.

The tolerance is stated in the same units of value as the variable in question; it is not a percentage. The tolerance is related to the value of the variable, not to expected value. Thus, entering a tolerance of 0.1 means that the actual location of any threshold will be within plus or minus 0.1. For example, if TreeAge Pro indicates finding a threshold at Var=0.391, this means that the threshold definitely occurs

somewhere between 0.381 and 0.401. Because TreeAge Pro applies linear interpolation after it meets your tolerance, you can expect the actual reported value to be even more accurate than the tolerance.

The Degree of non-linearity is used to try to avoid missing thresholds by dividing up the range into smaller intervals for the first pass. See note at the bottom of this section for more details.

The resulting Threshold Analysis Report is presented below.

Attribute	Variable	Var. Value	Strategy 1	Strategy 2	Exp. Value
EV	prob_extreme	0.167	Risky investment	CD Paying 5%	50

Threshold Analysis Report

The threshold analysis report shows the thresholds (rows) and information about each threshold (columns) in a table. The columns include:

- *Attribute*: The value for which thresholds were identified. This is usually EV for expected value.
- *Variable*: The variable for which the value was varied to identify thresholds.
- *Var. Value*: The value of that variable at the threshold.
- *Strategy 1, Strategy 2*: The strategies with equal expected value at that threshold.
- *Exp. Value*: The expected value of the two strategies at that threshold.



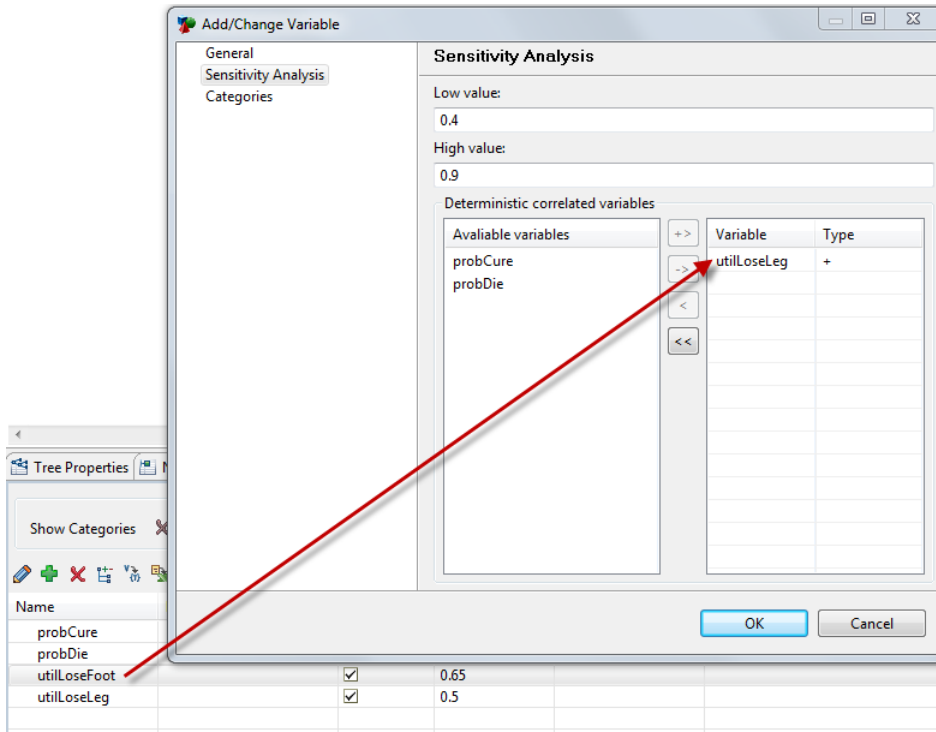
A sensitivity analysis may indicate multiple threshold values. However, this series of changes in policy will be identified correctly by TreeAge Pro only if the thresholds appear in different intervals in the first iterative pass. Since linear interpolation is used to find thresholds in a sensitivity analysis, only one threshold can be found per analysis interval.

For example, suppose that two thresholds exist in the same analysis interval, with optimality switching between the same decision options. Since the same policy is optimal at both ends, notwithstanding the intervening thresholds, TreeAge Pro will assume that no thresholds occur in that interval. There is no way to avoid this problem entirely. TreeAge Pro could subdivide a range into 100 intervals and still miss policy changes within an interval if the same optimal policy is specified at both ends. Even if different strategies are optimal at either end of an interval, and TreeAge Pro identifies a threshold in that interval, it is still possible that one or more additional thresholds in that same interval will have been missed.

For example, three alternatives, A, B, and C, might be compared using a sensitivity analysis; A is optimal at the beginning of an interval, B in the middle, and C at the end. Although you know that two thresholds (A to B, then B to C) actually occur, TreeAge Pro will find just one (a nonexistent one, A to C) from looking at the optimal alternative at the ends of the interval. The non-linearity hint is an attempt to minimize the likelihood this will occur. The more non-linear you describe the graph to be, the smaller the interval used by TreeAge Pro, so as to ensure catching any double thresholds.

18.6 Analyzing correlated variables

Linkages (i.e., perfect positive or negative correlations) between pairs of variables can be specified, for use during sensitivity analysis. These value linkages are set up in the Variable Properties View, Sensitivity Analysis Tab as described in the Working With Variables Chapter.



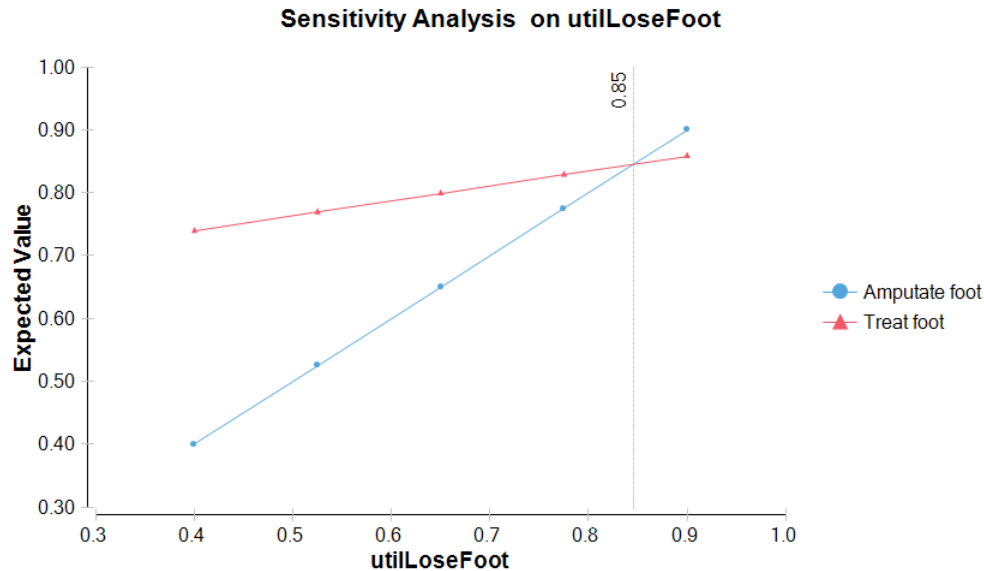
Correlated variables

Once created, the correlation is identified in the properties of both variables. Thus, when you choose to perform a sensitivity analysis on either member of a correlated pair of variables, TreeAge Pro will auto-adjust the correlated variable at the same time.

To perform a sensitivity analysis using correlated variables:

- Open the Special Features tutorial example tree, Correlated Variables. This tree already includes a positive correlation specified between the variables *utilLoseFoot* and *utilLoseLeg*. See above.
- Select the root, decision node.
- Choose Analysis > Sensitivity Analysis > One-Way... from the menu.
- In the Sensitivity Analysis dialog, select the variable *utilLoseFoot*. Specify 10 intervals and a range of 0.4 to 0.9. Click OK.

As the selected variable *utilLoseFoot* moves within its range (0.4 to 0.9), the correlated variable *utilLoseLeg* will move within its range (0.3 to 0.7).



Sensitivity Analysis with Correlated Variables

Note that the Treat Foot strategy's EV rises with the value of *utilLoseFoot* even though *utilLoseFoot* is not referenced by that strategy. This change is a result of the change in the correlated variable *utilLoseLeg*.

18.7 Additional sensitivity analysis topics

Other chapters cover additional software features which can be useful in testing the sensitivity of your trees, but whose use is not restricted to performing sensitivity analysis.

- Creating analysis sequences (Stored Analysis Abstracts and Sequences Chapter)
- Using variable sliders (Working With Variables Chapter)
- Linking trees (Tools and Functions for Complex Trees Chapter)
- Probabilistic sensitivity analysis (Probabilistic Sensitivity Analysis Chapter)

18.7.1 Sensitivity analysis on variables with non-numeric definitions

A sensitivity analysis can be performed on any variable in your tree, whether it has a numeric value definition (e.g., $X=1$ or $X=\text{Exp}(2)$) or a variable expression (e.g., $X=\text{Rate} \times \text{Util}$). When performing a sensitivity analysis on a variable defined as a formula, you have multiple options. You can perform a sensitivity analysis on the component variables (e.g., Rate and Util) using variable correlations or a multi-way sensitivity analysis. Alternatively, you can perform a one-way sensitivity analysis on the original variable (e.g., X) based on an estimated numeric value range.

If you treat X as the independent variable, however, the formula will be ignored during the course of the analysis. Definitions of Rate and/or Util at different points in your tree will not be used during this analysis.

It is advisable to focus sensitivity analysis on the finest-grain parameters. In the example above, X is no longer “finest-grain,” as it has been defined in terms of its two component variables. In general, models should be designed to ensure that the sensitive variables have a single numeric definition.

18.7.2 Checking probability coherence

Most forms of sensitivity analysis offer an option labeled Check coherence. When this option is selected, TreeAge Pro will ensure that, at each interval, (i) all probabilities sum to 1.0 and (ii) no probabilities are negative. The analysis will be halted if at any time either rule is violated.

If the subject variable is used to define a probability, you are encouraged to leave this option selected. This will ensure the validity of your model over the range of the analysis. This is particularly important in the initial stages of testing your model’s validity. The downside is that calculation time is increased. If calculation speed is a concern, and you are not including any probability variables in the sensitivity analysis, you may want to turn off coherence checking.

Also see the section on probability non-coherence in the Advanced Chance Node Techniques and Options Chapter.

18.7.3 Analyzing a single option

Normally, when performing a sensitivity analysis, a decision node is selected and TreeAge Pro displays one line for each of the alternative scenarios rooted at the selected node. It is possible to focus a one-way sensitivity analysis on a single scenario, rather than on all of the scenarios emanating from a decision node.

If the node you select prior to performing the sensitivity analysis is not a decision node, TreeAge Pro will assume that the results should be presented as a single line. This will represent the changing expected value of the scenario rooted at the selected node. (Healthcare module users: Note that this option is not available for cost-effectiveness sensitivity analyses, which must be performed at a decision node.)

If, however, you select a decision node which is an immediate descendant of a decision node, TreeAge Pro will give you the option of drawing one line for the selected node (as a branch of its parent), or multiple lines for the branches emanating from the selected decision node.

19. Monte Carlo Simulation, Distributions and Probabilistic Sensitivity Analysis

This chapter provides instructions on creating and referencing sampling distributions, performing Monte Carlo simulation, and interpreting the results.

The Distribution Functions, Options and Types Chapter provides details on functions for accessing distributions, options/customizations, and other usage details related to the built-in distributions available in TreeAge Pro. Instructions on creating custom, table-based distributions are found in the Creating and Using Tables Chapter.

For detailed information on using the Healthcare module to perform probabilistic sensitivity analysis and microsimulation on Markov and cost-effectiveness models, see the Cost-Effectiveness Simulation Reports and Graphs Chapters and the Individual-Level Simulation and Markov Models Chapter.

19.1 Uses of Monte Carlo simulation in modeling

The first section of this chapter summarizes some common applications of Monte Carlo techniques and tools in TreeAge Pro. The rest of the chapter focuses on setting up distributions in your model, and running probabilistic sensitivity analysis.

For tutorials on performing individual-level simulation (in particular, on Markov models), refer to the Individual-Level Simulation and Markov Models Chapter.

19.1.1 Monte Carlo overview

Each of the analyses described in previous chapters, including sensitivity analysis, is *deterministic*. There is no randomness in these types of expected value (EV) calculations: each parameter uses a specified point value, and every path through the model has a deterministic weight in the EV calculation based on its path probability (no matter how unlikely). If roll back or another analysis is repeated using the same parameters and tree, results do not change.

In contrast, there are many situations where it is useful to introduce random, or *stochastic*, elements into some part of the analysis. In such situations, Monte Carlo techniques can be applied.

A decision analysis model is a way of *visualizing* a potentially very complex equation. One level of uncertainty in a particular problem and its corresponding model is made visible with chance nodes: our uncertainty about the current and future state(s) of a modeled individual or experimental outcome. Discrete (or micro-) simulation can be used to explore this *1st-order uncertainty* (perhaps better labeled as *variability*). In decision analysis, the goal of such a simulation is generally still to calculate an expected value for each strategy being compared (in this case, an approximation based on long-run averaging of many random walks).

A separate, distinct level of uncertainty about a problem is, of course, *parameter uncertainty*. In this case, a parameter refers to some aspect of the system or world in which *all* our subjects, individuals, or experiments will operate (as opposed to some variable characteristic/state of a *single* subject). In some models and some systems, selected parameters (e.g., gravitational constant) may be known with precision, based on experimentation. In most decision analysis models, important system parameters (e.g., failure or survival rates) have not yet, or may never be, defined precisely through experimentation. Monte Carlo probabilistic sensitivity analysis (PSA) can be used to calculate an expected value over *2nd-order uncertainties* for a particular strategy.

Of course, in addition to reporting simple mean values, simulation “experiments” can be performed: individual-level, discrete simulation and/or probabilistic sensitivity analysis. Simulation output can be examined to account for uncertainty, calculating standard deviations/errors, percentiles or credible intervals, and a wide variety of other statistics.

Finally, multi-dimensional expectations can be carried out (when required), in which case the 2nd-order/parameter uncertainties (or value of information) are of primary interest and placed in an outer loop, while each repetition of the PSA expected value calculation may be approximated using a separate, inner, discrete simulation expectation/loop.

Additional overview of different Monte Carlo applications follows.



Discrete/microsimulation and probabilistic sensitivity analysis (PSA) have separate applications and require different interpretations. Note that commonly-used PSA outputs in health economics (e.g., value of information and acceptability curves, ICE scatterplot) may not have an intuitive application in a microsimulation. (In other words, the distribution of microsimulation outcomes is not interpreted the same way as the distribution of outputs from a PSA.)

19.1.2 Discrete (or micro-) simulation

Discrete simulation (microsimulation or 1st-order trials) is commonly used in the analysis of complex survival/failure models. Such simulation applications are covered in detail in the Individual-Level Simulation and Markov Models Chapter.

A much simpler, non-Markov discrete simulation, using the Stock Tree, is illustrated in the Analyzing Decision Trees Chapter. As described in the example, one way to look at a discrete simulation is as an approximation to expected value (EV) calculations in a decision tree. Simulation basically approximates EV calculations by sampling a representative distribution of paths through a model’s chance events. A discrete simulation of a complex model will often repeat as many “trials” as time allows, in order to improve the EV approximation.

In a single discrete simulation trial, a random walk (i.e., a series of uniform, pseudo-random numbers) selects a path through the chance nodes in the tree, with higher probability paths being more likely.

Running n simulation trials results in a list of n randomly-chosen outcomes, for example 10,000 “individuals” and their 20-year costs and life expectancy.

Although discrete simulation has less relevance in the average tree than in typical Markov projects, it is possible for very unwieldy trees to be greatly simplified or made more realistically by substituting probability distributions (e.g., Normal, Gamma, Exponential, etc.) for chance nodes, which are basically just discrete “distributions.” The required model changes are simple; the main sacrifice would be the requirement to use simulation instead of familiar EV analyses (like 1-way sensitivity analysis).

For example, the investment tree from the Analyzing Decision Trees Chapter could be revised with the risky investment chance node removed, and payoffs updated to include a likelihood distribution (e.g., Normal) representing change in investment value. Repeated investment decisions could also easily be modeled.

So, in discrete simulation models, variability can have two sources: chance node probabilities and parameterized distributions. In addition to possibly replacing event nodes in the tree, distributions have another, more common, application in simulation models. While simulation models sometimes assume a “homogenous cohort,” and include no initial variability in the characteristics of trials/entities, other models sample initial characteristics/states from probability distributions.

For example, a trial’s age might be sampled from a 1st-order distribution describing the population age distribution. Or, individuals might be bootstrapped from a table. (In an EV/cohort analysis model, in comparison, an initial series of chance nodes would explicitly separate subgroups based on patient characteristics, e.g., a few age groups.)

When parameterizing a distribution which represents 1st-order variability, simply change the sampling rate behavior from the default setting (for 2nd-order uncertainty) to instead sample per individual/1st-order trial.

19.1.3 Probabilistic sensitivity analysis

Like deterministic, n -way sensitivity analysis, Monte Carlo probabilistic sensitivity analysis (PSA) recalculates expected values in a tree multiple times, and is used to understand the impact of parameter uncertainties on the model results. One advantage of PSA is that any number of parameter uncertainties can be incorporated into an analysis. Sampling also enables greater weight to be placed on likely parameter values and combinations of parameters. PSA results estimate the total impact of uncertainty on the model, or the confidence that can be placed in the analysis results.

Probabilistic sensitivity analysis is covered in detail in the subsequent sections of this chapter.

19.1.4 Parameter distributions and model non-linearity

During a non-sampling analysis like rollback, all distributions in your model are fixed at their mean values. In some cases, taking an expected value from roll back will be equivalent to taking the sampling

mean from a probabilistic sensitivity analysis. However, this is not the case with all models and all parameters.

In a probabilistic sensitivity analysis on an uncertain rate in a survival model, for example, the sampling mean outcomes may differ significantly from the simple roll back “expected values” based on the point/mean value for the parameter. In such cases, the Monte Carlo PSA sampling means are the correct and preferred “expected values” for the model (assuming the likelihood distributions for the parameters are well-formed).



To support the complex, potentially lengthy types of analysis described in this chapter, TreeAge Pro can utilize up to eight processors on a single computer when performing Monte Carlo simulation.

19.1.5 EVPI/value of information analysis

Monte Carlo simulation can be used to perform various kinds of “value of information” analysis, similar to the structural form of EVPI described in the Analyzing Decision Trees Chapter.

The Analysis > Expected Value of Perfect Information command in TreeAge Pro calculates the difference between the baseline expected value of a decision, and the expected value when a chance node is temporarily shifted to the left of the decision. In a Monte Carlo simulation, the calculation of EVPI is done differently.

For example, if the optimal strategy changes depending on the values sampled for critical parameter uncertainties, then there would be some benefit to having “perfect information” about the uncertainty prior to the decision. The average of the values of each recalculation’s best option is the expected value with perfect information; it will either be equal to or greater than the best average value for any single alternative. Calculating the difference gives the expected value of perfect information.

Earlier versions of TreeAge Pro added EVPI reporting and charting options to the Monte Carlo simulation output window’s Graph popup menu. EVPI (and partial EVPI) are described later in this chapter.

19.2 Creating distributions

This section describes the basic steps for performing *probabilistic sensitivity analysis*:

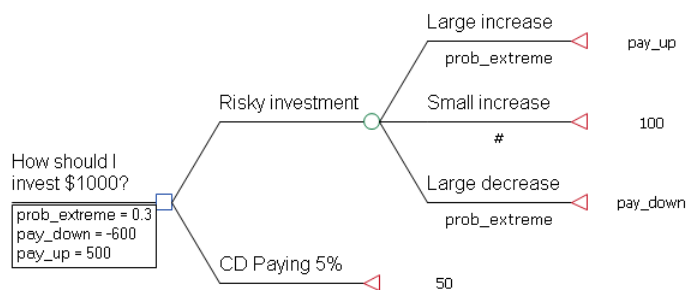
- *Define distributions* for your parameters;
- *Use the distributions* in the tree;
- *Run a Monte Carlo simulation*, to repeatedly: A) sample from distributions and B) recalculate expected values.

As detailed in the previous chapters on performing deterministic, *n*-way sensitivity analysis in TreeAge Pro, parameters targeted for analysis must be defined using *variables*. Similarly, before using Monte

Carlo simulation to perform probabilistic sensitivity analysis in TreeAge Pro, uncertain parameters must be defined using *distributions*.

Probability distributions can be employed in any formula in a tree, including variable definitions, payoffs/rewards, probabilities, and even parameters of other distributions. TreeAge Pro includes 20 built-in distribution types (Normal, Beta, etc.), as well as enabling sampling from custom, discrete “table” distributions. The Distribution Functions, Options and Types Chapter describes the functions and options available with distributions.

In TreeAge Pro, the distributions that you define are stored in a list in the tree like variables. Unlike variables, distributions are assigned an integer index (and a name, optionally). A distribution can be referenced in the tree with the *Dist(n)* function (where *n* is the distribution’s index) or alternatively by the distribution’s variable-type name. To illustrate the basic steps for probabilistic sensitivity analysis, the simple Stock Tree will be used. The tree – with variables but no distributions yet – is shown below.



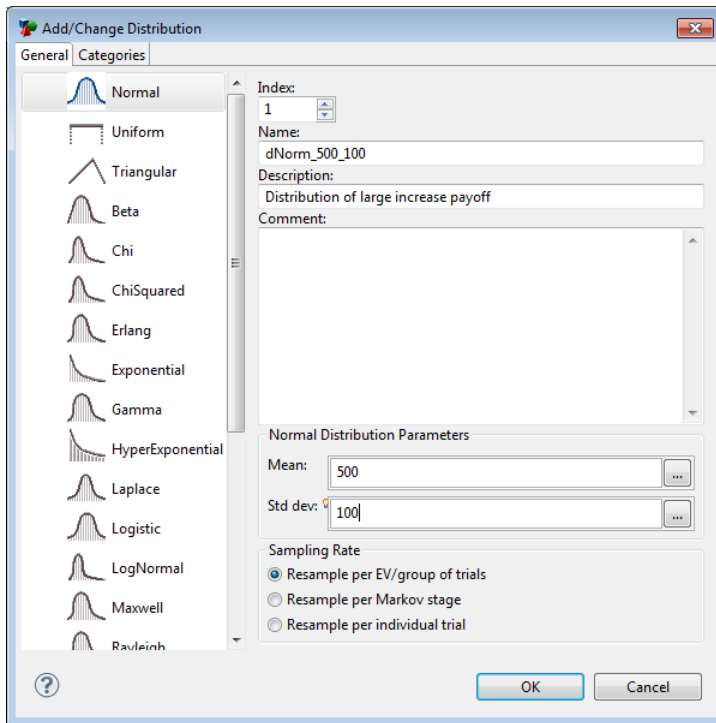
Three Variables Model

19.2.1 Defining a parameter using a distribution

The first steps are to change the payoff of the Large increase terminal node from a point estimate of 500 to a Normal distribution with a mean of 500 and a standard deviation of 100.

To define a distribution in the tree:

- Open the Get Started tutorial example tree "Three Variables".
- Save a copy named "Stock Simulation".
- Choose Views > Distributions from the menu.
- In the Distribution Properties View, click the "plus" toolbar icon to create a new distribution. The Add/Change Distribution dialog will open.
- Enter the new distribution's properties (type, name, description) and parameters (mean, std dev). The values are presented in the screen print below. After clicking OK, the new distribution will be listed in the Distribution Properties View (see below).



Add/Change Distribution Dialog



Use the Help icon in the Add/Edit Distributions Dialog to access the formulas TreeAge Pro uses for each distribution.

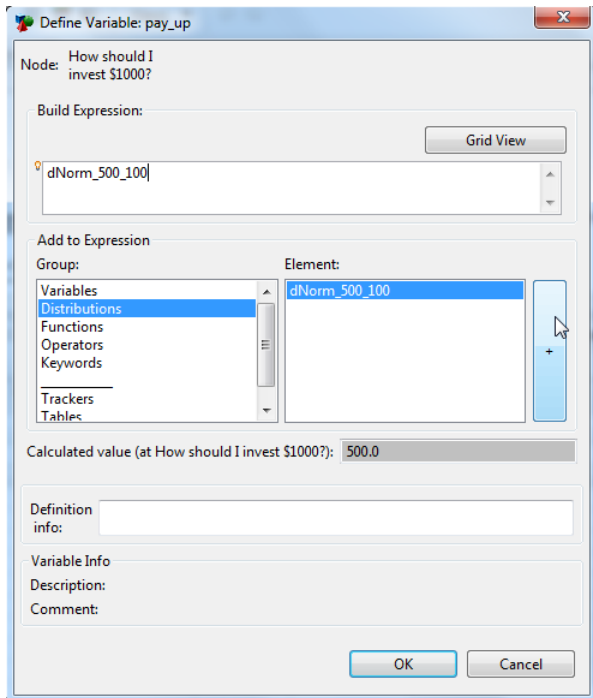
Variable Properties Distributions Tables Tree Properties Node Properties						
type filter text Clear						
Index	Type	Sampling Rate	Name	Description	Category	Resample at node: How should I i...
1	Normal	EV	dNorm_500_100	Distribution of large incr...		<input type="checkbox"/>

Distributions View

At this point, you have created the distribution, but it has not been integrated into the model. The new distribution represents the value for a high return from the risky investment. Therefore, it could be entered directly as the payoff value for that terminal node, or it could be used in the definition for the variable `pay_up`. We will use the second option.

To use the distribution in the `pay_up` variable definition:

- Right-click on the root node.
- Choose Define Variable > `pay_up` from the context menu.
- Clear out the existing definition (500).
- In the Group list, choose Distributions then double click on the distribution in the Element list.



Define pay_up with distribution dNorm_500_100

You could also have used the syntax *Dist(1)* instead of the distribution name since the distribution's index is 1.



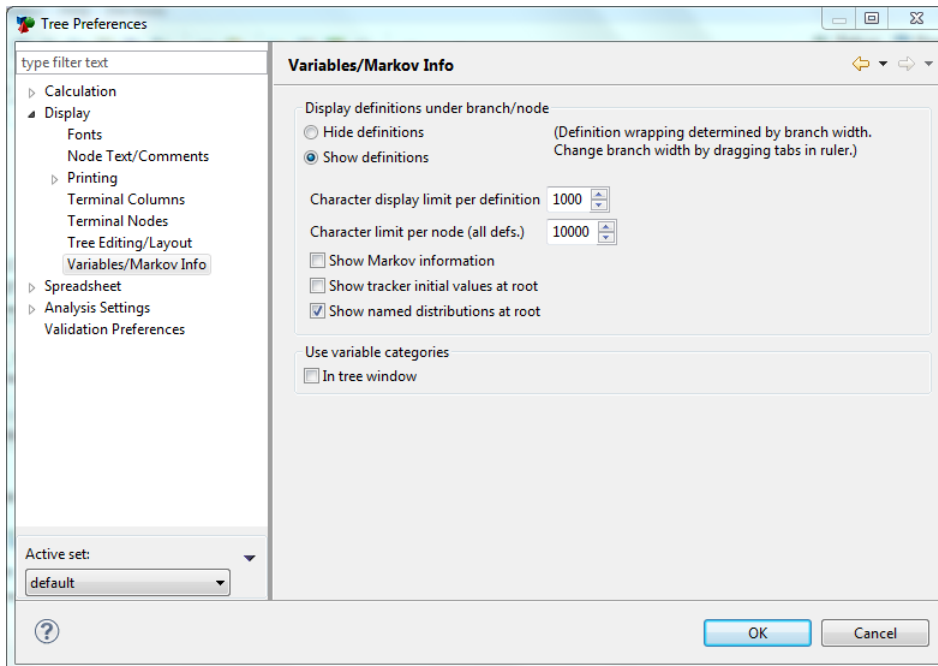
Distribution notes:

During roll back and other non-sampling analyses, a distribution is equal to its mean value. In the example, pay_up will still equal 500 during expected value calculations, because this was specified as the mean value of the Normal distribution.

The Dist() function in TreeAge Pro is equivalent to the DistSamp() function in DATA and earlier versions. Other versions of the distribution function, including DistForce() and DistKids(), are described in the next chapter.

During a single iteration of a sampling simulation, all references to a particular distribution return the same sample value because the default sampling rate (Once per EV) was used. The sampling rate can be set to resample more frequently, for example in Markov microsimulations.

Each named distribution can be presented in a list at the root node of the tree. To show the list on the face of the tree, turn on the "Show named distributions" option within the Display > Variables\Markov Info category of Tree Preferences.



Tree Preferences - Show named distributions

19.2.2 Distributions View

Distributions are managed (added, edited and deleted) through the Distributions View. This is a tree-level view since distributions are "global" within the context of a tree (like variable properties, but not like variable definitions).

Below is a screen print of the view showing one of each of the possible distribution types supported by TreeAge Pro.

Index	Type	Name	Description	Category
1	Normal	Distribution1	Normal	
2	Uniform	Distribution2	Uniform	
3	Triangular	Distribution3	Triangular	
4	Beta	Distribution4	Beta	
5	Chi	Distribution5	Chi	
6	ChiSquared	Distribution6	Chi-Square	
7	Erlang	Distribution7	Erlang	
8	Exponential	Distribution8	Exponential	
9	Gamma	Distribution9	Gamma	
10	HyperExponent...	Distribution10	Hyper-Exp	
11	Laplace	Distribution11	Laplace	
12	Logistic	Distribution12	Logistic	
13	LogNormal	Distribution13	Log-Normal	
14	Maxwell	Distribution14	Maxwell	
15	Rayleigh	Distribution15	Rayleigh	
16	Weibull	Distribution16	Weibull	
17	Poisson	Distribution17	Poisson	
18	Binomial	Distribution18	Binomial	
19	Fractile	Distribution19	10-50-90	

Distributions View

The toolbar executes the following functions within this view.

- *Edit*: Open the Add/Change Distribution Dialog to edit the properties and parameters of a single distribution selected in the list.
- *Add*: Add a new distribution via the Add/Change Distribution Dialog.
- *Add new distributions*: Add multiple distributions via the Add/Change Distribution Dialog.
- *Delete*: Delete the distribution(s) selected in the list.
- *Group by Category*: Show the distributions list grouped by category.
- *Edit in Excel*: Users with the optional Excel Module can output the distribution properties and parameters to Excel, where they can be edited and returned to TreeAge Pro.
- *Report*: Generate a report showing parameters and EV values for each distribution.
- *Graph It*: Sample from the distribution and create a graph displaying the sampled values.
- *Highlight*: Highlight the distribution within the model in the Tree Diagram Editor.

Distributions View Functions

The next chapter contains additional information on managing distributions.



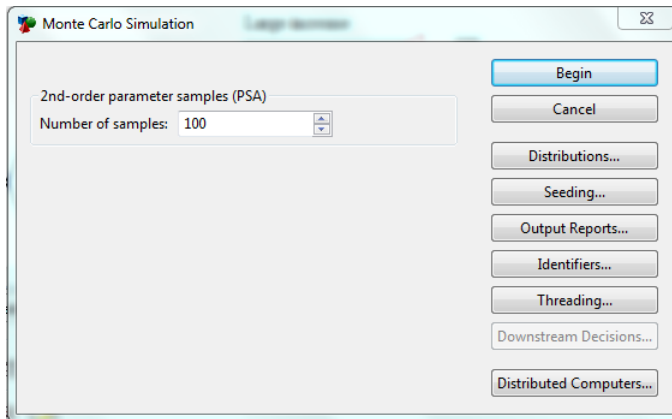
If your license includes the Excel module, the Edit in Excel button can be used to export selected distributions to a table in a new Excel worksheet where they can be edited. From Excel, the Add or Update Distributions command in the TreeAge Add-In menu can be used to update the active tree with any changes. Refer to the Graphing, Reporting and Modeling Using Excel Chapter for details.

19.3 Performing probabilistic sensitivity analysis

After defining and using distributions in your tree, you can run a Monte Carlo simulation to see how resampling parameter values affects calculations at a selected node. Simulations can be run at any node except a terminal node, making it possible to analyze only part of a tree.

To perform a probabilistic sensitivity analysis/Monte Carlo simulation:

- Select the root node of the "Stock Simulation" tree built in the prior section.
- Choose Analysis > Monte Carlo Simulation > Sampling (Probabilistic Sensitivity)...from the menu. The Monte Carlos Simulation Dialog will open.



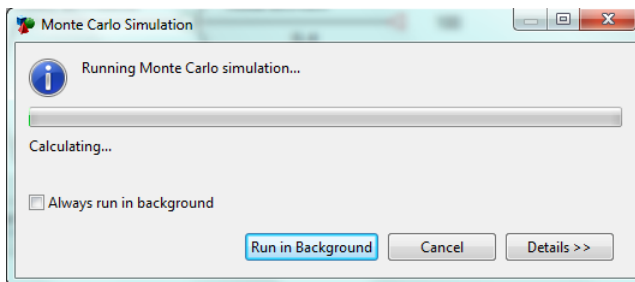
Monte Carlo Simulation Dialog

If you have defined distributions in the Distributions View (*even if they are not used anywhere in the tree*), the Monte Carlo Simulation Dialog will present a variety of options for the analysis. In this case, you simply specify the desired number of distribution samples and corresponding model recalculations.

Now, specify the settings to use for this simulation:

- Change the number of distribution samples to 500.
- Click Begin to start the simulation.

In running this simulation, TreeAge Pro will randomly sample 500 values from the distribution defined in the tree and recalculate expected values for the branches of the selected decision node based on each parameter sample.



Monte Carlo Simulation Progress Dialog

The simulation output window shows the progress of the simulation. During long running simulations, it may be helpful to run the analysis in the background to allow you to continue working in TreeAge Pro. You can also click the Cancel button to stop the analysis.

Once the simulation is complete, a final statistical summary will be displayed. The summary includes mean, standard deviation, and other summary statistics for each strategy.

Statistic	Risky investment	CD Paying 5%
Mean	9.498	50.000
Std Deviation	31.191	0.000
Minimum	-72.286	50.000
2.5%	-50.624	50.000
10%	-33.108	50.000
Median	13.050	50.000
90%	49.156	50.000
97.5%	71.220	50.000
Maximum	75.328	50.000
Sum	949.814	5,000.000
Size (n)	100.000	100.000
Variance	972.907	0.000
Variance/Size	9.729	0.000
SQRT[Variance/Size]	3.119	0.000

Probabilistic Sensitivity Analysis Output

If a simulation is performed at a decision node, the summary statistics for each branch are presented from left to right starting with the top branch. The statistics report can be exported to several formats (HTML, Excel, Word, etc.) by clicking the "Export Report As..." toolbar icon in the simulation output window. The statistics report can also be viewed within TreeAge Pro in HTML or flat grid format by clicking the appropriate links under the heading "Report Options...". The statistics report can also be exported to files in those formats via the "Export Report As..." toolbar icon.

Note that the mean value above for the *Risky investment* strategy is close to but is not equal to 10 (the expected value without simulation). If more iterations (samples) were selected for the simulation, that mean would move closer to the expected value. Note that there is no variance in the *CD Paying 5%* strategy since it does not reference a distribution.



Since there is only one symmetrical distribution (normal) in the model, the mean from the simulation should approach the mean from expected value analyses. However, the mean from simulations will not always approach the mean from expected value analyses as noted in the non-linearity section.



Unless you use "seeding," there should be statistical variation from one simulation to the next.

19.3.1 Saving Monte Carlo simulations

Depending on the complexity of the model, and the number of samples and recalculations you specify, running a simulation can be time consuming. For this reason, the simulation output window can be saved separately from the tree as a report output (*.rptx) file. Saving the simulation output will allow

you to share the complete results with other TreeAge Pro users, or to generate graphs and reports from the simulation at a later time.

To save the Monte Carlo output window:

- Select the Monte Carlo output as the active view.
- Choose File > Save As... from the menu or click the Save toolbar icon in the simulation output window.
- Select the appropriate project, enter a file name and click OK.
- The *.rptx file will be saved in the project.

You will notice the extension *.rptx will be added to the title of the tab displaying the simulation output. You can open the *.rptx file later to reopen the output file.



Monte Carlo simulation output files grow relative to the number of rows and columns. Sizes of ~1 GB would be achieved with 1 million rows and 150 output columns. Use a Zip compression program to shrink file size.



Output *.rptx files contain database tables holding the report output data. Refer to the Technical Details Chapter for information on querying the data using an external tool.

19.3.2 Probability coherence

Sampled probabilities can sometimes generate a set of branch probabilities that do not sum to 100%. This will cause the PSA simulation to fail. There are a few options to correct the non-coherent probabilities:

1. Use beta distributions for probabilities; Beta distributions always generate samples between 0 and 1.
2. Use Dirichlet distributions for chance nodes with more than two branches. A single Dirichlet distribution will generate all branch probabilities, guaranteeing that the sum is 100%.
3. Normalize probabilities to force non-coherent probabilities to sum to 100%.

Correcting non-coherent probabilities

19.3.3 Basic simulation reports and graphs

Clicking the *Strategy Values* link shows the calculated expected value for each sampled iteration of the simulation (see below). Note that each iteration generated a different value for the *Risky investment* strategy (which referenced distributions), but the same value for the *CD Paying 5%* strategy (which did not reference distributions).

Monte Carlo Strategy Values

Showing page 1 of 20

Sample	Risky investment	CD Paying 5%
1	8.3119950803	50
2	59.2081301434	50
3	-15.2524209776	50
4	-27.358643051	50
5	-17.2702291635	50
6	78.7662410185	50
7	-23.5377638583	50
8	-7.8847914839	50
9	45.2500370549	50
10	10.2707576494	50
11	23.0952940026	50
12	-45.9047800107	50
13	-0.6427684884	50
14	-8.8100188434	50
15	-4.3165391282	50
16	36.5386713487	50
17	61.2871904352	50
18	-2.1403219194	50
19	-36.3943194346	50

PSA Output - Strategy Values

The Strategy Values report window can handle thousands of rows, but it's broken down into pages for viewing. This output can also be exported to external files as can most reports and graphs.

The *Values*, *Dists*, *Trackers* link displays complete output from each iteration of the simulation, including:

- *Strategy values*: The EV for each strategy (assuming the simulation was run at a decision node).
The Strategy Key relates the column headings back to the strategies' node labels.
- *Input distributions*: The value of each input distribution.
- *Tracker values*: The final value of each tracker (for each Microsimulation trial).

Monte Carlo Simulation Report

ITERATION	STRATEGY_1	STRATEGY_2	DIST_1
1	8.3119950803	50	494.3733169344
2	59.2081301434	50	664.027100478
3	-15.2524209776	50	415.8252634079
4	-27.358643051	50	375.4711898299
5	-17.2702291635	50	409.0992361218
6	78.7662410185	50	729.220803395
7	-23.5377638583	50	388.2074538056
8	-7.8847914839	50	440.3840283871
9	45.2500370549	50	617.5001235163
10	10.2707576494	50	500.9025254982
11	23.0952940026	50	543.6509800087
12	-45.9047800107	50	313.6507332976
13	-0.6427684884	50	464.5241050388
14	-8.8100188434	50	437.2999371887
15	-4.3165391282	50	452.2782029061
16	36.5386713487	50	588.4622378289
17	61.2871904352	50	670.9573014508
18	-2.1403219194	50	459.5322602685
19	-36.3943194346	50	345.3522685515
20	-23.0320282915	50	389.8932390284
21	24.9985403601	50	549.9951345337

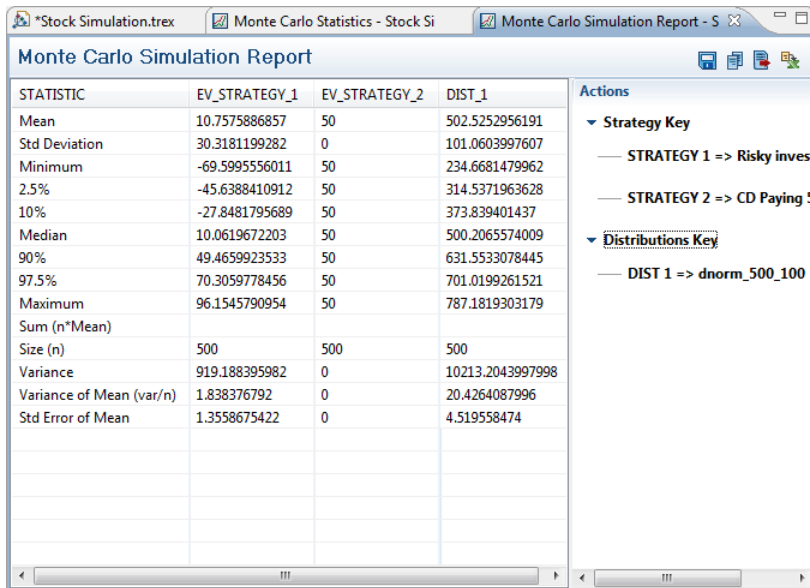
Actions

- Strategy Key
 - STRATEGY 1 => Risky investment
 - STRATEGY 2 => CD Paying 5%
- Distributions Key
 - DIST 1 => dnorm_500_100

PSA Output - Values, Dists, Trackers

The *Values, Dists, Trackers (Tab-Delimited XLS/TXT)* link creates a report which will export directly to *.csv/*.txt. This is much more efficient for large data output than running the Values, Dists, Trackers report first, then exporting that to Excel.

The *Statistics - Values, Dists, Trackers* link displays a summarized output from the simulation. The data columns are the same as for the "Values, Dists, Trackers" link, except that statistical information is presented.



The screenshot shows a web browser window with three tabs: "*Stock Simulation.trex", "Monte Carlo Statistics - Stock Si", and "Monte Carlo Simulation Report - S". The active tab displays the "Monte Carlo Simulation Report". The report contains a table with statistical data for three variables: EV_STRATEGY_1, EV_STRATEGY_2, and DIST_1. The table includes rows for Mean, Std Deviation, Minimum, 2.5%, 10%, Median, 90%, 97.5%, Maximum, Sum (n*Mean), Size (n), Variance, Variance of Mean (var/n), and Std Error of Mean. To the right of the table is an "Actions" panel with expandable sections for "Strategy Key" and "Distributions Key".

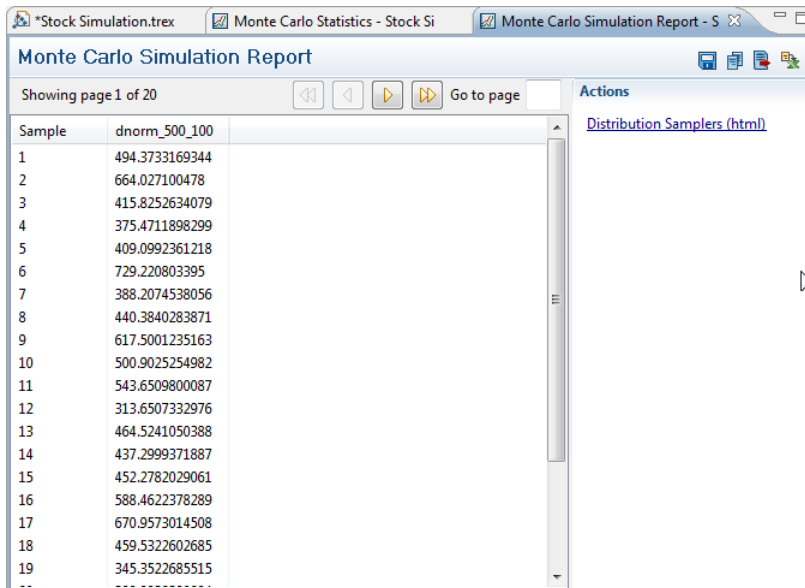
STATISTIC	EV_STRATEGY_1	EV_STRATEGY_2	DIST_1
Mean	10.7575886857	50	502.5252956191
Std Deviation	30.3181199282	0	101.0603997607
Minimum	-69.5995556011	50	234.6681479962
2.5%	-45.6388410912	50	314.5371963628
10%	-27.8481795689	50	373.839401437
Median	10.0619672203	50	500.2065574009
90%	49.4659923533	50	631.5533078445
97.5%	70.3059778456	50	701.0199261521
Maximum	96.1545790954	50	787.1819303179
Sum (n*Mean)			
Size (n)	500	500	500
Variance	919.188395982	0	10213.2043997998
Variance of Mean (var/n)	1.838376792	0	20.4264087996
Std Error of Mean	1.3558675422	0	4.519558474

Actions

- ▼ **Strategy Key**
 - STRATEGY 1 => Risky invest
 - STRATEGY 2 => CD Paying 5
- ▼ **Distributions Key**
 - DIST 1 => dnorm_500_100

PSA Output - Statistics - Values, Dists, Trackers

The *Sampling Distributions* link displays the distribution samples associated with each iteration of the simulation. If there were more than one distribution in the model, additional links would be available. The distribution output can be displayed in an HTML format as well.



The screenshot shows a web browser window with three tabs: "*Stock Simulation.trex", "Monte Carlo Statistics - Stock Si", and "Monte Carlo Simulation Report - S". The active tab is "Monte Carlo Simulation Report - S". The page title is "Monte Carlo Simulation Report". Below the title, it says "Showing page 1 of 20". There are navigation buttons (back, forward, search, etc.) and a "Go to page" input field. On the right, there is an "Actions" section with a link "Distribution Samplers (html)". The main content area is a table with two columns: "Sample" and "dnorm_500_100". The table contains 19 rows of data, with the first row being "1" and "494.3733169344".

Sample	dnorm_500_100
1	494.3733169344
2	664.027100478
3	415.8252634079
4	375.4711898299
5	409.0992361218
6	729.220803395
7	388.2074538056
8	440.3840283871
9	617.5001235163
10	500.9025254982
11	543.6509800087
12	313.6507332976
13	464.5241050388
14	437.2999371887
15	452.2782029061
16	588.4622378289
17	670.9573014508
18	459.5322602685
19	345.3522685515

PSA Output - Sampling Distributions

For all simulations, probability distribution histograms can be displayed for the model output calculations, as well as for any sampling distributions. Other graphs are limited to specific types of simulations (i.e., sampling, trials, etc.) and characteristics of the model itself (i.e., cost-effectiveness vs. simple calculation method).

Under the heading "Charts", the following graphs and groups are listed:

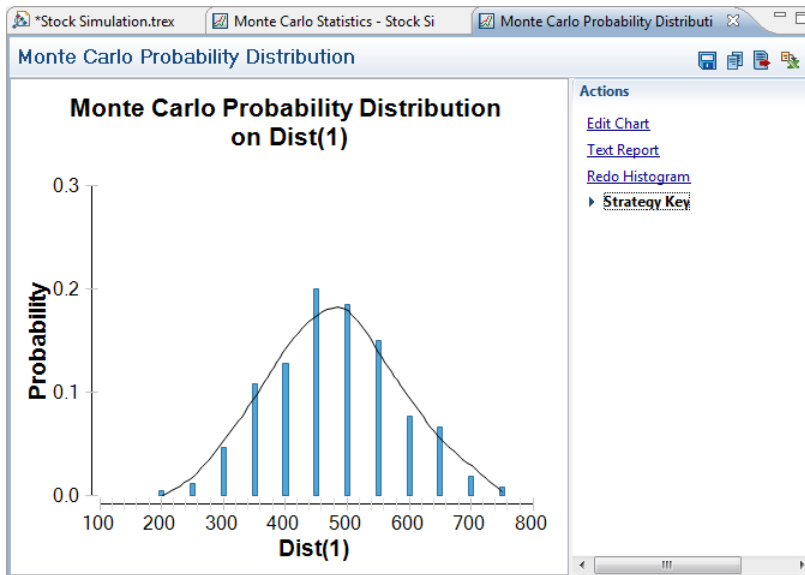
- *Sampling Distributions*: A set of probability distributions for the samples from each input distribution.
- *Strategy Selection Frequency*: A probability distribution for which strategy was selected as optimal.
- *Output Distributions*: A set of probability distributions for each expected value and tracker output.
- *Other Charts*: Other output options associated with specific types of models and analyses.

Chart group options from Monte Carlo output



Cost-effectiveness probabilistic sensitivity analysis generates additional output. These outputs are described in the Cost-Effectiveness Simulation Reports and Graphs Chapter.

The *Sampling Distributions* group allows you to generate a probability distribution graph for each of the distributions sampled in the analysis. In this case, there is only one - Dist(1). When you click on a link for a distribution, you are prompted for an approximate number of bars to include in the probability distribution. After entering a value, the probability distribution is displayed.



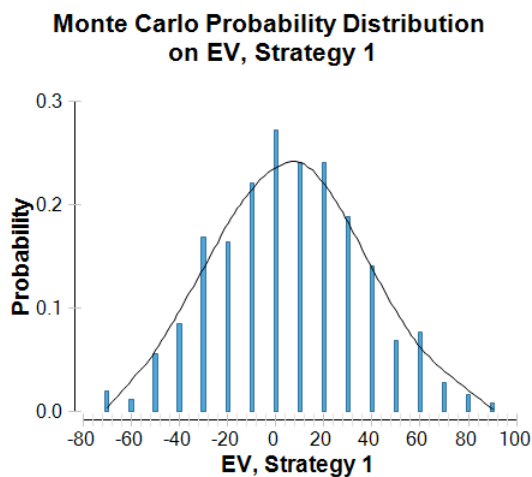
Parameter Distribution



Note that the Redo Histogram link allows you to regenerate the graph with more or fewer bars. For a discussion of the options available in TreeAge Pro for working with and customizing distribution graphs, refer to the Graphs Chapter.

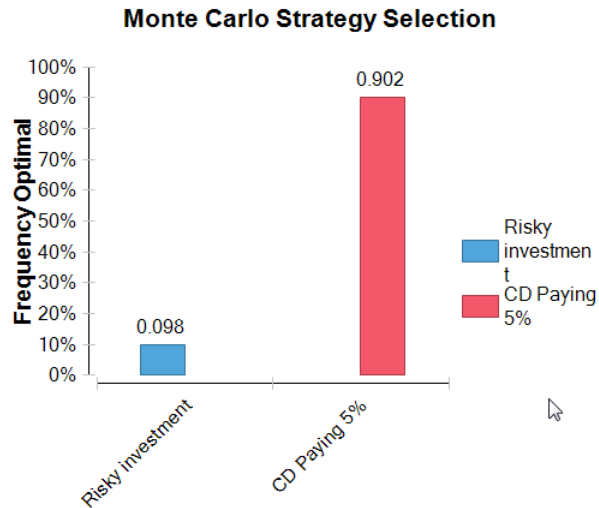
The *Output Distributions* group generates probability distributions for outputs from the model. In this case, there is a single subgroup - *EV* (expected value) with an item for each of the two strategies. Other models/simulations can include other types of outputs.

The graph below shows the probability distribution for the Risky Investment strategy's expected value.



Output Distribution for Risky investment strategy

The *Strategy Selection Frequency* graph shows the samples for which each alternative is optimal. You will be prompted for an indifference threshold which specifies the minimum significant difference between strategies such that one is optimal. The graph is presented below.



Strategy selection graph



If you have the TreeAge Pro add-in for Excel, graphs generated as TreeAge graph files can also be created as Excel charts. In some cases, Excel charts have different/additional functionality. Refer to the Graphing, Reporting and Modeling Using Excel Chapter for details. Numerous additional graphs and reports are available in cost-effectiveness simulations performed using the Healthcare module for TreeAge Pro. For example, in a cost-effectiveness simulation, the Acceptability Curve is used in place of the Strategy Selection graph. Refer to the Cost-Effectiveness Simulation Reports and Graphs Chapter for additional details.

19.3.4 Expected Value of Perfect Information (EVPI/EVPPI) reports and charts

Probabilistic sensitivity analysis simulations performed at decision nodes include a detailed EVPI report and summary chart. Both the report and chart are included under the "Charts" heading of the simulation output.



The EVPI reported for a microsimulation (versus a sampling simulation) might be interpreted as the value of being able to predict all chance nodes' outcomes. Using the Stock Tree, for example, compare the Analysis > Expected Values of Perfect Information result at the single chance node in the model to the simulation EVPI report/chart in a simulation performed at the decision node.

The calculation of EVPI in simulations is relatively straightforward:

1. determine the *overall optimal strategy*, using the simulation mean values (for cost-effectiveness models, uses net monetary benefits based on a specified threshold ICER);
2. determine the *optimal strategy for each sample iteration* in the simulation (normally the outermost loop will be a sampling loop, although TreeAge Pro will do similar calculations even if it is just a microsimulation with no sampling);

3. for each iteration, if the optimal strategy is not the overall optimal strategy, calculate its *incremental value* (iteration optimal - overall optimal), which will be ≥ 0 ;
4. report the *average (expected) value of perfect information* over all iterations.

Steps for EVPI calculation in simulation

The simulation run earlier in this chapter generates an *EVPI\EVPPi Summary Report* link. Clicking on this link generates the following report.

EVPI	Optimal Strategy
1.582	2

EVPI/EVPPi Summary Report

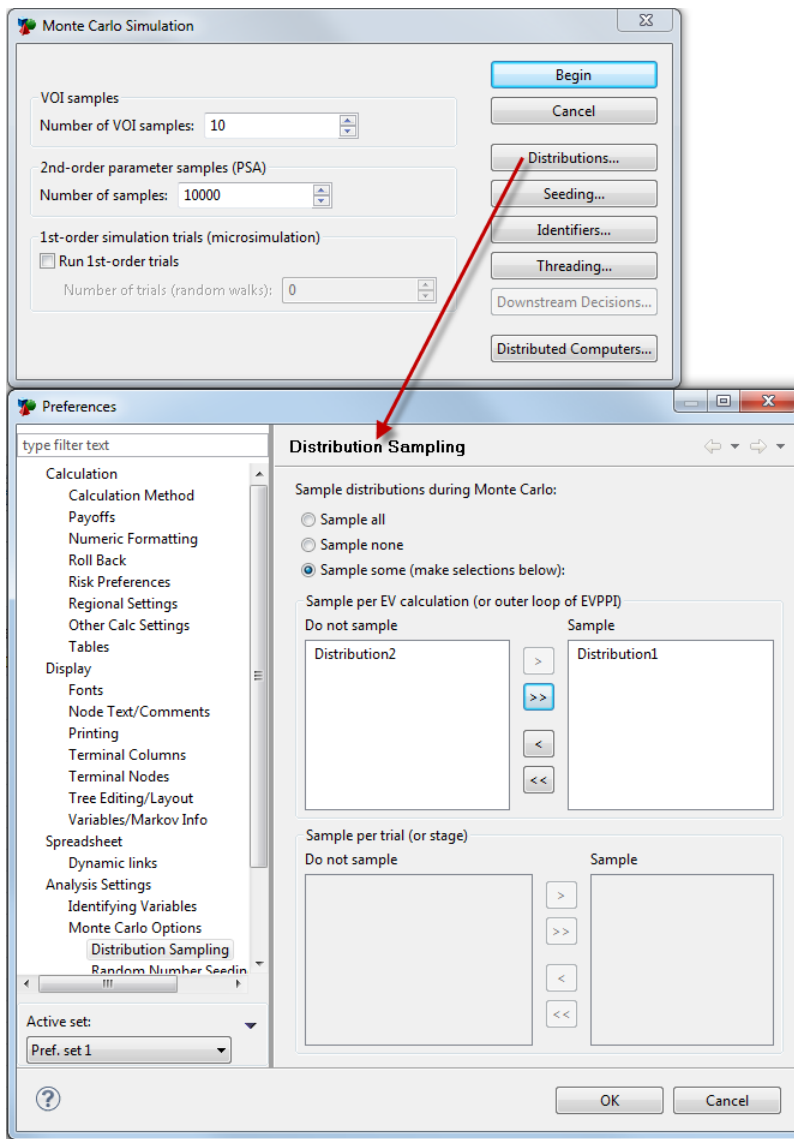
The EVPI value reflects the total EVPI from each iteration divided by the number of iterations as described above. To see the details from the individual iterations, click the EVPI\EVPPi Details link.

ITERATION	EVPI	ITEROPTIMAL	ITEROPTIMALEV	OVERALLOPTIMAL	OVERALLOPTIMALEV
1	0	2	50	2	50
2	9.2081301434	1	59.2081301434	2	50
3	0	2	50	2	50
4	0	2	50	2	50
5	0	2	50	2	50
6	28.7662410185	1	78.7662410185	2	50
7	0	2	50	2	50
8	0	2	50	2	50
9	0	2	50	2	50
10	0	2	50	2	50
11	0	2	50	2	50
12	0	2	50	2	50
13	0	2	50	2	50
14	0	2	50	2	50
15	0	2	50	2	50

EVPI/EVPPi Details Report

To perform partial EVPI (or EVPPi), usually a two-level sampling loop is required. To handle this analysis robustly, in case of any nonlinearity in model parameters, TreeAge provides a 3-dimensional simulation option (3rd dimension can be a microsimulation loop).

In an EVPPi two-level sampling loop, use the Distribution Sampling options to select the distributions to sample in the outer sampling loop. The remaining distributions will be sampled in the inner sampling loop. In the example below, Distribution 1 is sampled in the outer loop and Distribution 2 is sampled in the inner loop.



Monte Carlo Distribution Sampling

The parameter(s) of interest are sampled in the outer sampling loop N times; for each outer iteration, an inner loop samples the remaining uncertainties (and recalculates expected values) M times, in case of non-linearities in these remaining distributions.

For more information on EVPI/EVPPI simulation options in TreeAge Pro, refer to the current recommendations on performing EVPPI and related analyses found in the journal of the Society for Medical Decision Making (and elsewhere):

<http://www.smdm.org/>

19.4 Simulation options

This section describes options you can select when running simulations.

19.4.1 Multi-threading

To better handle lengthy simulations, TreeAge Pro will use multiple processors on a single computer when performing Monte Carlo simulation.

If you are running simulations on a dual-processor computer, you can instead specify to use only one simulation thread, which will leave one processor idle during the simulation (allowing other programs to run quickly while the simulation is running). If you are running a multi-threaded simulation, it is recommended that you leave the Optimize all expressions... setting on.



It is sometimes useful to run a simulation with a single thread to make it easier to read the debugging output. Otherwise, output from multiple threads can get mixed together.

19.4.2 Distributed simulations

TreeAge Pro provides a mechanism to run Monte Carlo simulations on multiple computers. This can help speed up long simulations. Distributed "slave" computers are setup in Application Preferences for use with any model.

To setup distributed computers:

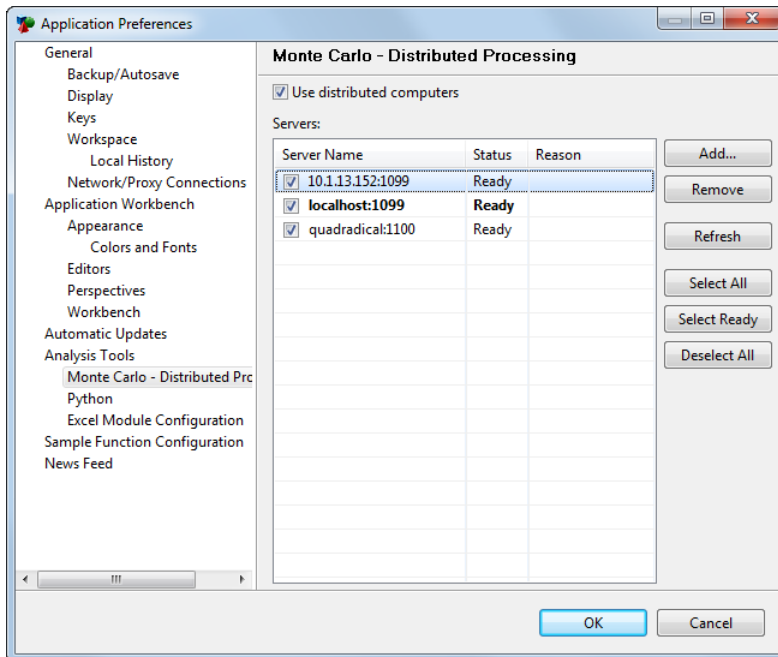
- Choose Window > Application Preferences from the menu.
- Select the category Analysis Tools > Monte Carlo - Distributed Processing
- Check the box labeled Use distributed computers.
- Click the Add button and enter the slave's computer name or IP address. Click OK.
- Check the boxes next to the slaves you want to use for the simulation.
- Click OK to save the Application Preferences changes.

Setup distributed computers



You can enter an IP range to scan for available distributed computers on your network. For example, enter 192.168.1.100-192.168.1.200 to search for distributed computers within that IP address range.

Once a set of slave computers is added, the list will look like the image below.



Application Prefs - Distributed Computers

**Distributed Simulation Licensing/Software Versions:**

- Master computers must have active Maintenance to use distributed slave computers.
- Slave computers require no TreeAge Pro license.
- Master and slave computers must be running the identical software build version. If the versions do not match, the slave computer will not be used for the analysis. You can check the version in the Application Preferences.

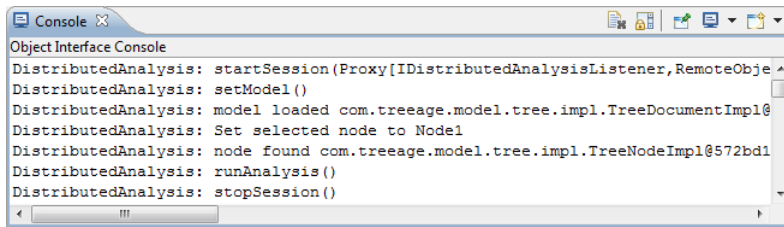
To use slave computers, the "Use distributed computers" box must be selected and individual computers must be selected. *The master computer must be selected in the list of Distributed Computers in order to use it for simulation batches.* This allows you to choose whether to utilize the master computer's system resources.

When you then run a Monte Carlo simulation, the master computer will send batches of iterations to each slave computer. The slave computers will return the iterations back to the master computer, where they are collected and eventually reported back through the user interface.



Note that there is overhead associated with passing data back and forth among computers, so you will not see the simulation speed double by adding a second identical computer for iteration batches.

On the slave computers, you can see that batches are being processed within the Object Interface Console. See below.



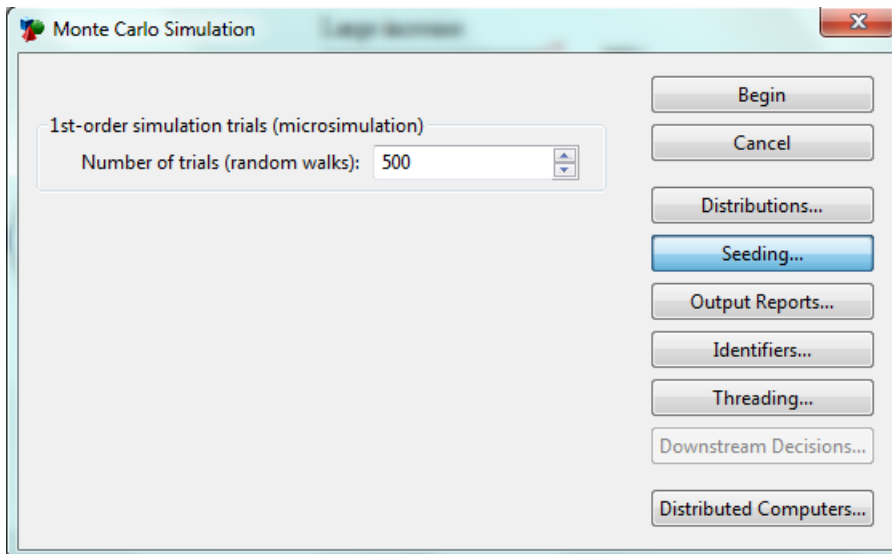
Object Interface Console - Distributed Analysis Output

19.4.3 “Seeding” the random number generator

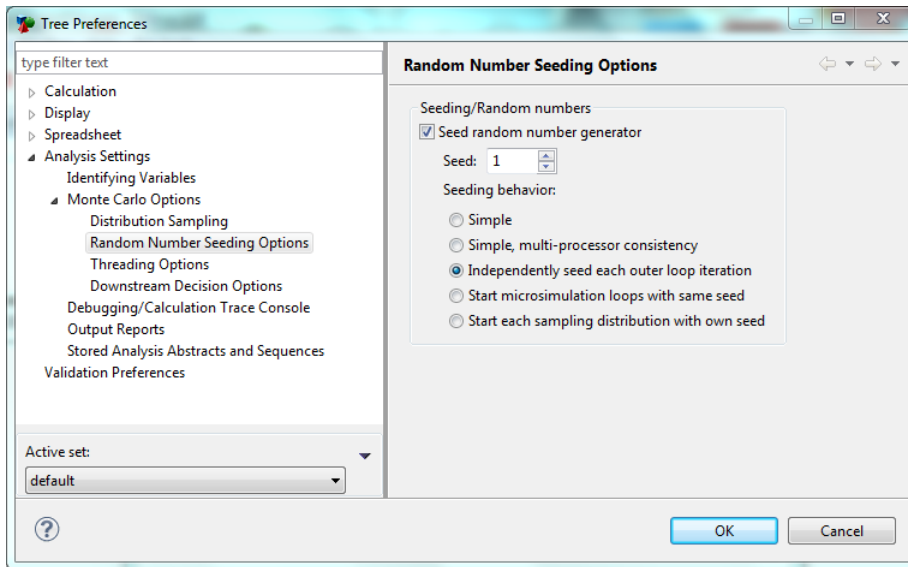
Normally, Monte Carlo simulations use dynamic information from the computer’s clock to initialize a sequence of pseudo-random numbers. If you ever need to force the same set of samples and/or trials for several different simulations of the same model, you can specify to use the same, predictable sequence of pseudo-random numbers.

To seed a simulation:

- In the setup dialog for a Monte Carlo simulation, click the Seeding... button. This opens the Preferences Dialog to the Analysis Settings > Monte Carlo Options > Random Number Seeding Options Category.
- Check the Seed random number generator box, and specify an integer seed value (from 1 to 64,000).
- Select a seeding behavior.
- Click OK to save the Tree Preferences.



Monte Carlo Simulation Setup - Seeding



Seeding Tree Preferences

The seeding behavior options are described below.

- *Simple*: uses a repeatable sequence of numbers.
- *Simple, multi-processor consistency*: ensures that the sequence of random numbers does not change depending on the number of processors used.
- *Independently seed each outer loop iteration*: generates a sequence that does not change if strategies are reordered, removed, or excluded.
- *Start microsimulation loops with same seed*: uses consistent seeds for inner loops in a multidimensional simulation. This reduces variance among inner loop iterations.
- *Start each sampling distribution with own seed*: provides for better consistency in distribution sampling in EVPPi (i.e., 2-level sampling) simulations. Without this setting, rerunning a simulation after moving a single distribution from the outer sampling loop to the inner loop (or vice versa) could cause all other, unmoved distributions to nonetheless return new series of sample values. ONLY USE FOR EVPPi SIMULATIONS.

Seeding behavior options



The seeding options (and several other simulation options) are stored with the model's Tree Preferences. The preference values are saved when you save the model.

All seeding should be used cautiously, and not as a substitute for running a representative number of iterations.

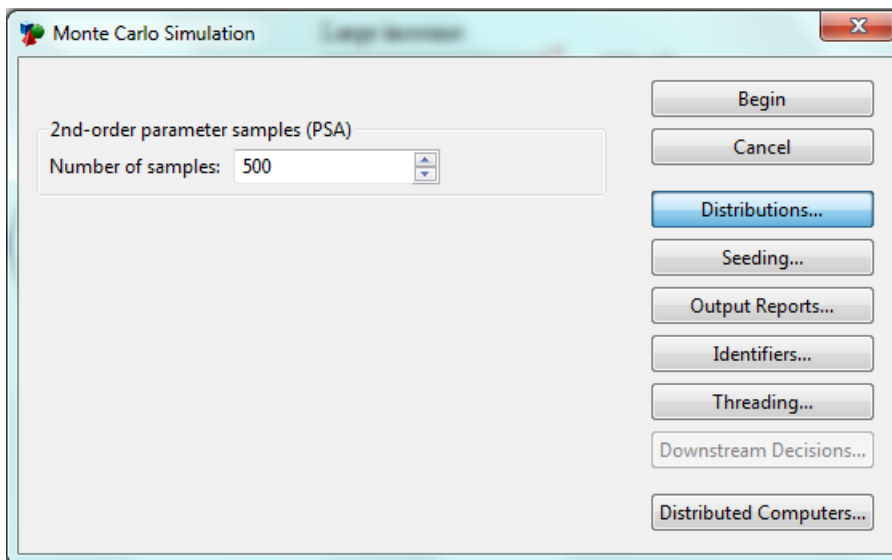
In special situations, it may be useful to utilize the $\text{Seed}(n)$ function, which overrides the current position of the random number generator (in the currently executing simulation thread). Different values of n will result in different subsequent random number sequences.

19.4.4 Turning off sampling of selected distributions

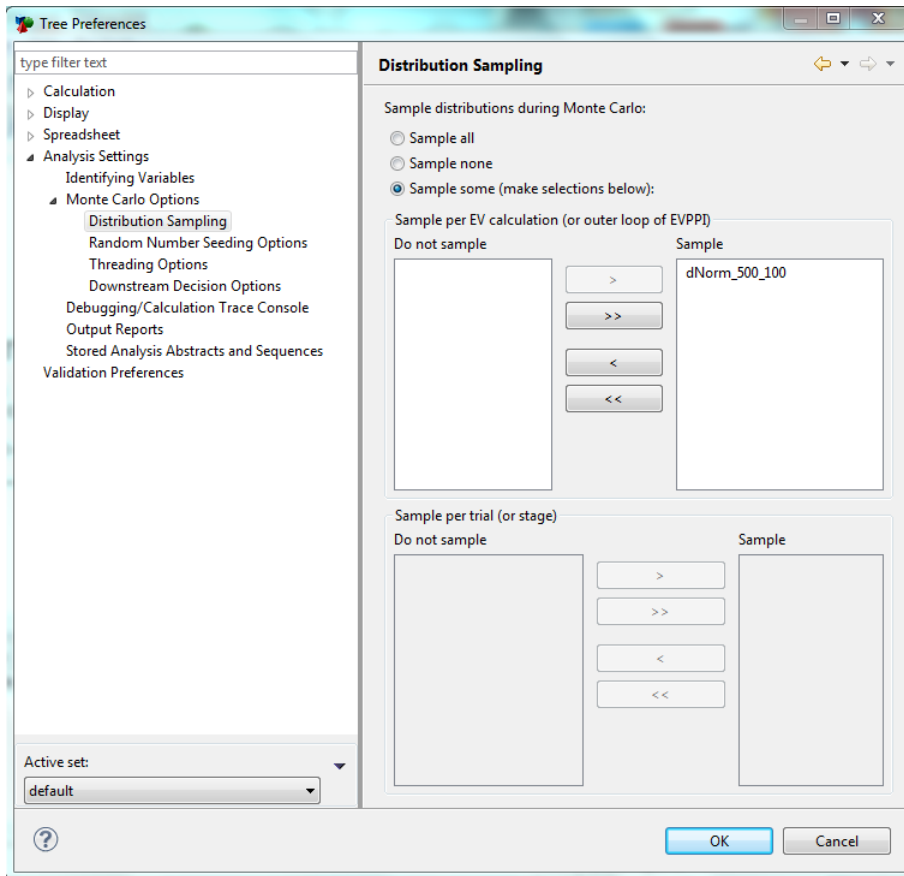
You have the option of specifying that only a subset of the distributions should be sampled during a simulation. Non-sampled distributions will be set at their mean.

To sample from selected distributions during a simulation:

- In the setup dialog for a Monte Carlo simulation, click the Distributions... button. This opens the Tree Preferences Dialog to the Analysis Settings > Monte Carlo Options > Distribution Sampling Category.
- Select a "Sample which distributions" option.
- If the option "Sample some" is selected, use the left/right buttons to move the appropriate distributions between the available list on the left to the sample list on the right.
- Click OK to save the Tree Preferences.



Monte Carlo Simulation Setup - Distributions



Distribution Sampling Tree Preferences

19.4.5 Identifying simulations

It can be useful to associate identifying information with simulation output, especially when you have more than one simulation output window open at once, or if you are saving and reopening or sharing the Monte Carlo simulation files. Simulation output can be "labeled" with *identifying variable* values and with a *simulation comment*.

Identifying variables

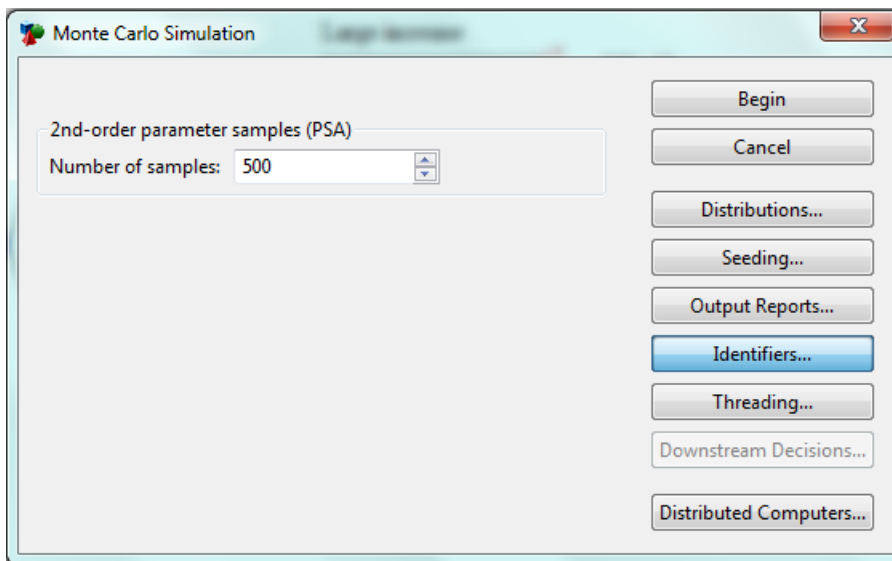
Simulation output windows can be labeled using the values of variables in your tree. The resulting simulation output window will display a list of the variables and their default values.

Monte Carlo simulation identifying values are particularly useful in combination with the stored analysis features in TreeAge Pro (Stored Analysis Abstracts and Sequences Chapter).

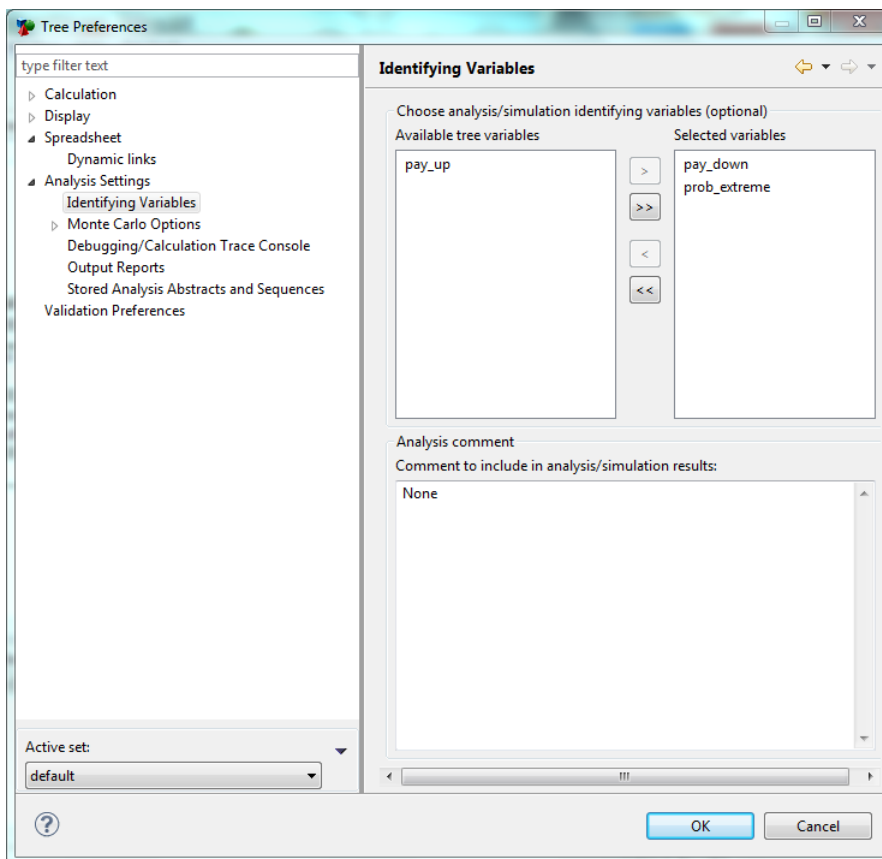
To add identifying variable values in the simulation window:

- In the setup dialog for a Monte Carlo simulation, click the Identifiers... button. This opens the Tree Preferences Dialog to the Analysis Settings > Monte Carlo Options > Identifying Variables Category.

- Use the left/right buttons to move the appropriate variables between the available list on the left to the identifier list on the right.
- Click OK to save the Tree Preferences.



Monte Carlo Simulation Setup - Identifiers



Identifying Variables Tree Preferences

Note that the pay_up variable was not selected because its value is defined using a distribution, and those distribution values would already be included in the simulation output.

The Identifying Variables are stored in the Tree Preferences, so that you do not need to re-select them for each new simulation.

After the simulation is executed, the values of the identifying variables are available through the simulation output's "Identifying Variables" link.

Monte Carlo Simulation Identifying Variables

<u>VARIABLE</u>	<u>VALUE</u>	<u>NOTES</u>
pay_down	-600	
prob_extreme	0.3	

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Monte Carlo simulation output - Identifying Variables

19.4.6 "Downstream" decision nodes during microsimulations

If your tree includes decision nodes to the right of the node where a simulation is being performed, each trial must select a single path when it encounters such embedded decisions. For simulations using only expected value calculations (e.g., PSA with no trials), TreeAge Pro always re-evaluates the optimal path at the embedded decision, and utilizes the optimal strategy's expected value.

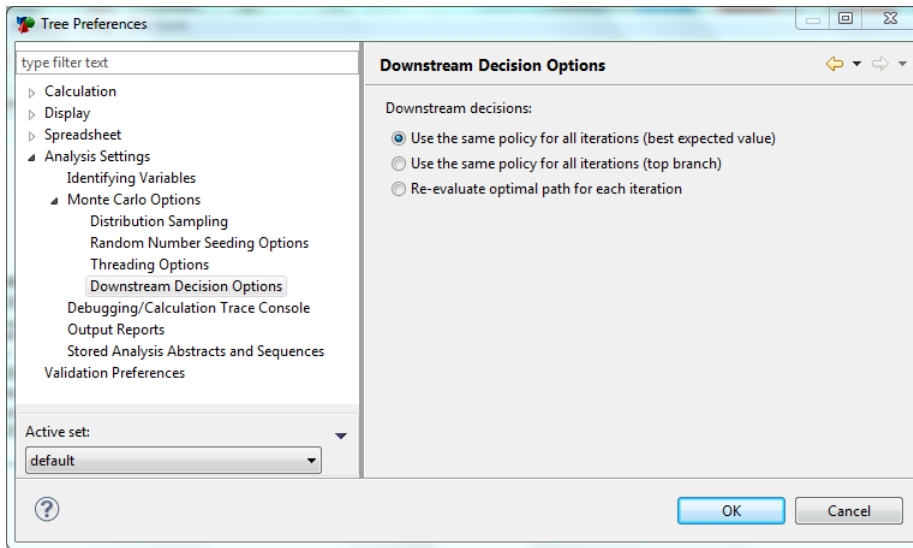
For trials, however, embedded decision nodes can be handled in three ways:

1. based on expected values calculated prior to the simulation, a single optimal policy can be followed for all iterations;
2. select the topmost branch;
3. if any parameter uncertainty (e.g., PSA) distributions are used in the tree, the optimal policy can be re-evaluated for each sample iteration based on the current distribution sample values.

Downstream Decision Options

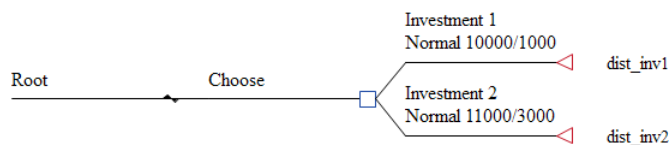
To set the downstream decision algorithm for a simulation:

- Open the Tree Preferences dialog.
- Choose the category Analysis Settings > Monte Carlo Options > Downstream Decision Options.
- Select the option that is appropriate for your model.
- Click OK to save the Tree Preferences.



Tree Preferences - Downstream Decision Options

The Special Features tutorial example model Downstream Decisions illustrates the three options.



Downstream Decisions tree

Note that the root node is not a decision node, so the Choose node is a downstream decision node. Therefore, the downstream decision options selection will control which investment is selected for each trial.

The payoffs for the two investments are both distributions sampled by EV/set of trials. The investment 2 option has a larger mean value, but the two distributions' standard deviations are large enough that some PSA iterations will favor investment 1.

If the "Use the same policy for all iterations (best expected value)" option is selected, then all trials select investment 2 because it has a larger mean value.

ITERATION	STRATEGY_1	DIST_1	DIST_2
1	10627.0238952042	9310.0238559047	10627.0238952042
2	10627.0238952042	9310.0238559047	10627.0238952042
3	11537.6391462628	11387.3507894617	11537.6391462628
4	11406.2566360602	10851.103077771	11406.2566360602
5	11070.5295069834	11338.203498946	11070.5295069834
6	8184.721863118	8748.8651350691	8184.721863118
7	15728.7660168421	9912.3821302521	15728.7660168421
8	8098.2265790949	11272.426283851	8098.2265790949
9	16115.8718751078	8422.5999651573	16115.8718751078
10	7162.5670091455	8908.7424801344	7162.5670091455

Downstream decisions simulation - option 1

If the "Use the same policy for all iterations (top branch)" option is selected, then all trials select investment 1.

ITERATION	STRATEGY_1	DIST_1	DIST_2
1	9455.8420048169	9455.8420048169	8963.8534217272
2	9179.5724291185	9179.5724291185	12714.3602673479
3	11053.4817380021	11053.4817380021	16066.05323632
4	8490.4066274874	8490.4066274874	8813.8027660347
5	7965.4507344791	7965.4507344791	11621.2058105319
6	8901.2750243677	8901.2750243677	7867.7306662844
7	10396.3096842637	10396.3096842637	11654.662505989
8	11094.8643512052	11094.8643512052	7811.0323825064
9	10886.8393147013	10886.8393147013	11646.9940127368
10	9103.9889099047	9103.9889099047	6163.3291320817

Downstream decisions simulation - option 2

If the "Re-evaluate optimal path for each iteration" option is selected, then all trials will select the investment with the larger sampled value. Note that in iterations 8 and 10, the trials selected investment 1 rather than investment 2.

ITERATION	STRATEGY_1	DIST_1	DIST_2
1	12799.1518411715	9789.5802963149	12799.1518411715
2	12799.1518411715	9789.5802963149	12799.1518411715
3	12581.2113116797	10941.4309918236	12581.2113116797
4	12110.0991938226	7494.9585645685	12110.0991938226
5	15472.8330733108	10294.9335975723	15472.8330733108
6	15759.9991641418	12515.4115334148	15759.9991641418
7	12925.617297642	10555.5148503688	12925.617297642
8	10872.2151270027	10872.2151270027	10501.4004689819
9	13995.4929073821	10012.7439080292	13995.4929073821
10	9420.9358688952	9420.9358688952	8314.1662492857

Downstream decisions simulation - option 3

19.4.7 Distribution options

Each distribution created in a tree can be extensively customized. Refer to the Distribution Functions, Options and Types Chapter for complete details on distribution types, functions, and sampling behavior options.

19.5 Customizing simulations

The previous section described the various simulation options that can be explicitly set prior to running the simulation. For more complex situations, there are additional methods at the modeler's disposal for customizing the behavior of simulations.

19.5.1 The TreeAgeProLib (Excel add-in)

The TreeAgeProLib library enables writing macros to control setup and execution of simulations via the Object Interface. Refer to the Using the TreeAge Pro Object Interface Chapter for more details.

19.5.2 Keywords

A variety of simulation keywords (built-in counters) can be used in expressions (e.g., tracker modifications, rewards, probabilities) during simulation. The table below describes each keyword.

Keyword	Description
<code>_trial</code>	Index for the trial/iteration being processed in the microsimulation loop
<code>_sample</code>	Index for the sample/iteration being processed in the sampling/PSA loop
<code>_voi_sample</code>	Index for the VOI iteration being processed (the outer most simulation loop in EVPI/EVPPI analysis)
<code>_trial_size</code>	Number of trials/iterations for the microsimulation loop
<code>_sample_size</code>	Number of samples/iterations for the PSA simulation loop
<code>_voi_sample_size</code>	Number of iterations for the outer most simulation loop in EVPI/EVPPI analysis
<code>_state_index</code>	Index for the Markov state counting down from the top state (which is 1)
<code>_thread_index</code>	Index for the thread which is running all or a portion of the overall simulation (simulations can run as multiple threads to increase the speed of the simulation)
<code>_parallel_trials_clock</code>	The current <code>_clock</code> value during a parallel trials discrete event simulation that uses the <code>_CLOCK</code> value to keep the trials synchronized.
<code>_parallel_trial_creator</code>	If you turn off probability coherence, you can create individuals in a parallel trials simulation. This keyword can be used by a new trial to get the number of the trial that created it (i.e., via a transition/chance event with non-coherent probs).
<code>_parallel_trials_set</code>	The index for the current set of parallel trials.
<code>_parallel_trials_sets_size</code>	The number of sets of parallel trials in the simulation.
<code>_cache_level</code>	When using the <code>Node()</code> function to access parts of the tree, this returns the "depth" of the calls from nested <code>Node()</code> functions.

Simulation keywords

The most useful will be: `_trial` and `_sample`. These allow you to reference the trial index and/or sample index in expressions within the tree.



For information on the parallel trials keywords, refer to the Individual-Level Simulation and Markov Models Chapter.

19.5.3 Monte Carlo special variables

Normally, the only calculations that are done between iterations of a microsimulation or probabilistic sensitivity analysis are distribution resampling. However, user-defined expressions can also be

evaluated at strategic points between simulation iterations. Expressions assigned to the following regular variable names will be evaluated.

Special Variable	Expression is executed...
<code>_monte_pre_trial_eval</code>	Before each trial is run
<code>_monte_post_trial_eval</code>	After each trial is run
<code>_monte_pre_sample_eval</code>	Before each sample is run.
<code>_monte_post_sample_eval</code>	After each sample is run.
<code>_monte_pre_info_sample_eval</code>	Before each VOI sample is run.
<code>_monte_post_info_sample_eval</code>	After each VOI sample is run.
<code>_post_analysis</code>	After entire analysis is complete.

Monte Carlo Special Variables

If you create a variable with one of the above names and define it at the root node, it will be evaluated at the appropriate time during the analysis (see above). This can be used to perform additional reporting functions, such as recording to global matrix individual inner loop results occurring within a two-dimensional simulation.

For example:

```
_monte_post_sample_eval = GlobalN(_voi_sample;_sample;Node(1;0;1;1))
```

would record the expected value for the top arm of the simulated node.



These keywords can be used to initialize and output global matrices at specific times during the simulation. For example, you can output trial-level data within a two-dimensional simulation where the standard simulation output would only provide mean values for the trial sets.

Use the `_post_analysis` keyword to output data when the entire analysis is complete. Note that the `_post_analysis` keyword can be run for other analyses, while the other keywords are only executed within the context of Monte Carlo simulation.

19.6 Two- and three-dimensional simulations

In most models, a probabilistic sensitivity analysis should recalculate expected values for each new set of parameter samples to see the effects of parameter uncertainty.

However, there is an alternative method for recalculating the tree after each set of parameter samples: estimating an expected value by averaging many first-order trials. This process – estimating expected values by averaging a sufficiently large number of random, individual outcomes – is sometimes called microsimulation.

19.6.1 Recalculate using first-order trials (microsimulation)

In most models, two-dimensional simulation for the purposes of probabilistic sensitivity analysis is unnecessary. It is normally preferable to run a 1-dimensional loop that recalculates EVs for each set of randomly sampled parameter values – except in the case of certain types of models, for example:

1. Markov models using *tracker variables to follow detailed event history*. Refer to the Individual-Level Simulation and Markov Models Chapter for details on trackers.
2. Trees using *distributions to represent variability among individuals* (much like a chance node does) rather than just parameters whose values are uncertain. Refer to the Distribution Functions, Options and Types Chapter for more information on sampling rates.

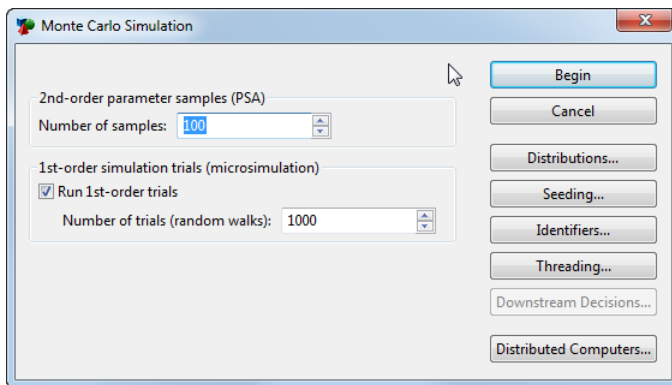
Scenarios requiring two-dimensional simulation for PSA

Both of these situations call for microsimulation (described at the beginning of this chapter).

Depending on the complexity of the model, *each microsimulation* may require thousands of first-order trials in order to adequately *approximate an expected value*. Therefore, in the context of a probabilistic sensitivity analysis, using a microsimulation for each recalculation of the tree (for a new set of samples) can be very time consuming. This is why two-dimensional simulation is generally avoided unless a model requires microsimulation to calculate an expectation.

To perform probabilistic sensitivity analysis using first-order microsimulation trials for model recalculations:

- Select the root node of your tree, or a node to the right, to set the context for analysis.
- Choose Analysis > Monte Carlo Simulation > Sampling + Trials... from the menu.
- Enter the number of samples for the 2nd-order parameter loop.
- Enter the number of trials for the 1st-order trials loop.



Monte Carlo Simulation setup for two-dimensional simulation



Specify a sufficient number of 1st-order trials to get a good average for each new sample value (or set of values). Initially you should experiment with smaller number of iterations in both the sampling loop as well as the inner, microsimulation loop.

In the context of Markov models using trackers with microsimulation, refer to the Individual-Level Simulation and Markov Models Chapter for more information on two-dimensional simulations.

19.6.2 Two-dimensional simulation details

In microsimulation models also requiring probabilistic sensitivity analysis, a two-dimensional (or two-loop) approach to is used:

1. N number of parameter samples and I number of microsimulation trials are specified.
2. Draw one set of samples for the parameter distributions.
3. Holding the sampled parameter values constant, a group of I microsimulation trials are performed. The mean of the results from the trials is reported (e.g., cost and/or effectiveness values, as well as tracker values).
4. Steps 2 and 3 are repeated for each of the N sets of parameter samples.

Two-dimensional simulation detailed steps

Once the simulation is complete, the final report will include N rows, showing each set of parameter samples and the corresponding *mean values* for I microsimulations. The distribution of these results reflects the total uncertainty resulting from the parameter distributions.

Note that the detailed results for each of the I trials within each sample are not reported.

19.6.3 Three-dimensional, value of information simulations

To perform partial EVPI (or EVPPI), usually a two-level sampling loop is required. TreeAge provides a 3-dimensional simulation option to handle these more complex values of information simulations. In this case, the innermost loop or “dimension” allows for a set of microsimulation/individual trials to be used in the innermost expectation step of the EVPPI simulation (if required).

Obviously, as noted in the previous section, multi-level/multi-dimensional Monte Carlo simulations can be very time consuming. Distributed simulation can divide the required processing time among multiple computers. And, if carefully used, special microsimulation error reduction seeding options can be used to reduce, to some extent, the number of sampling iterations required to get stable 2nd-order uncertainty measures.

20. Distribution Functions, Options and Types

This chapter provides information about each of the built-in sampling distribution types available in TreeAge Pro.

Derivations and detailed explanations of the distribution formulae provided here may be found on many math/statistics web sites, and in most texts on probability theory. See, for example, Christensen, Ronald; Data Distributions: A Statistical Handbook (2nd Ed.); Lincoln, Massachusetts: Entropy Limited, 1989.

Refer to the following chapters for information on using distributions in trees.

- More Sensitivity Analysis Tools
- Monte Carlo Simulation, Distributions and Probabilistic Sensitivity Analysis
- Advanced Chance Node Techniques and Options
- Individual-Level Simulation and Markov Models Chapter

20.1 Distribution functions

Distributions can be referenced in two different ways: by name (if an optional name is defined in the distribution properties) or by index. The following functions employ the argument index to reference a distribution defined for the tree. With the exception of DistKids(), they are intended primarily for use during Monte Carlo simulation.

Function	Description
Dist(index)	During expected value calculations, this function returns the indexed distribution's mean. During Monte Carlo simulations, this function returns the most current random sample for the distribution (unless the distribution is flagged to not sample).
Dist(index; column)	For a multi-variate distribution (i.e., a Dirichlet, Multivariate Normal or multi-column table distribution), returns the sample value (or mean, if not sampling) from the designated value column.
Dist(index; 1)	For a univariate distribution (i.e., not Dirichlet, Multivariate Normal or multi-column table), forces a sample from the distribution, even during non-sampling analyses. It is recommended that you use the DistForce function rather than this function (see below). For a table distribution, use the first value column for probabilities.
Dist(index; 2)	For a univariate distribution (i.e., not Dirichlet, Multivariate Normal or multi-column table), returns the stored sample value for a virtual branch created using the DistKids() syntax, below. For a table distribution, use the second value column for probabilities.
DistForce(index)	Unlike the Dist(index) syntax, DistForce() always samples a value, even during non-sampling analyses such as roll back.

Function	Description
DistForce(index; column)	DistForce(index; column) also works for multi-variate distributions. Note that DistForce(index; 1) must be called first in order to trigger a new set of sample values for all columns; then the stored sample values for the other columns can be referenced.
DistKids(index; samples)	A probability function which dynamically creates invisible branches (number = samples) at a chance node during any calculation. Used in combination with the Dist(index; 2) syntax described above.
DistProb(index; val)	For the specified distribution, returns the approximate cumulative probability of the specified value.
DistTrim(index; min; max)	Same as the Dist() function, except it will resample (up to a maximum of 10 times) until a sample is returned that falls between the specified minimum and maximum values.
DistValue(index; prob)	For the specified distribution, returns the approximate value at the specified cumulative probability.
Seed(n)	Sets the random number generator in the current thread (multi-processor simulations use multiple threads). Use a positive integer argument up to 2 billion. If no argument is given, or a zero argument, a pseudo-random seed is generated (i.e., as if no seeding).
DistSamp(index)	Obsolete. Use the Dist() function.

Distribution Functions



The DistTrim will usually generate a value between the min and max arguments. However, it is possible for 10 consecutive samples to fall outside this range. This function can be used in conjunction with the Min and Max functions to guarantee a value within the range.

Min(maxValue; Max(minValue; DistTrim(distIndex; minValue; maxValue)))

20.2 Distribution options

Some special options can be used to make distributions more flexible.

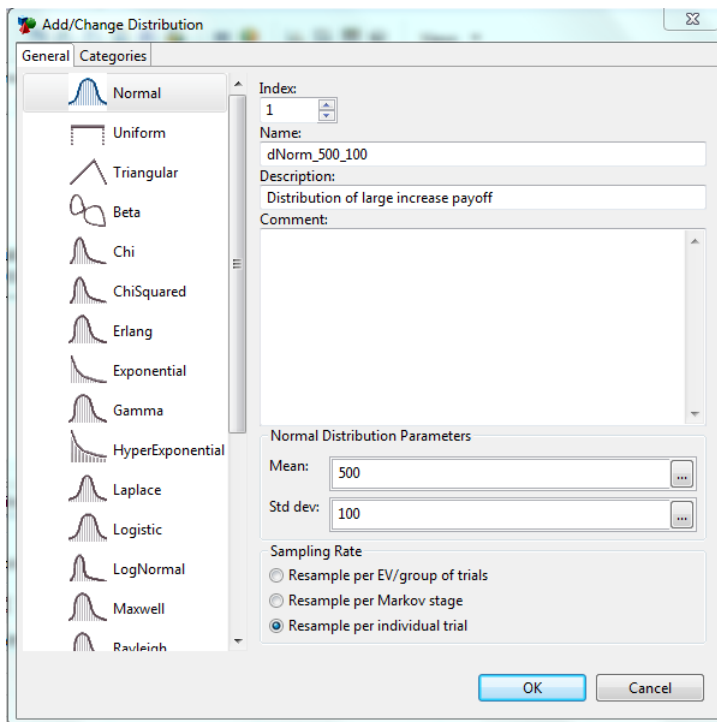
20.2.1 Changing sampling rate for microsimulation models

While most distributions are used to sample possible values of uncertain parameters for probabilistic sensitivity analysis, distributions can also be used to instead represent individual variability/patient characteristics. These two different classes of distributions should be identified via the TreeAge Pro distribution's sampling rate property. This is particularly important if a model includes both types of distributions.

To change the sampling frequency for a particular distribution:

- Open a model with at least one distribution.
- Choose Views > Distributions from the toolbar.

- Select a distribution from the list.
- Click the "edit" toolbar button to open the Add/Change Distribution dialog.
- Select the Sampling rate option "Once per individual trial".



Add/Change Distribution Dialog - set sampling rate

This option will be of interest in specific models:

1. Markov models using tracker variables to follow detailed event history, including discrete event models.
2. Trees using some distributions to represent variability among individuals (in the same way that a chance node could).

Characteristics of models that might use distributions that sample once per trial

By default, distributions are set to sample once per tree EV recalculation. In a two-dimensional simulation, this would be per "group of trials". By default, a simple microsimulation resamples only distributions set to sample per 1st-order trial or Markov stage.

It is also possible to set a distribution's sampling frequency to generate a new sample value at each Markov cycle/stage during first-order trials and/or cohort/EV calculations. The `DistForce()` syntax can be used to resample more frequently.

20.2.2 Sampling during non-simulation, EV calculations

During non-simulation analyses, such as roll back and sensitivity analysis, a reference to a distribution normally returns the *mean* value every time it is referenced. The `Dist()` function can, however, override

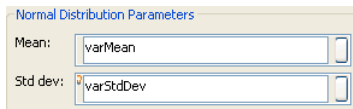
this behavior and return a randomly sampled value from the referenced distribution during any EV calculation.

To cause a random distribution sample to be returned by the Dist() function during expected value (EV) calculations, simply add a second parameter to the function with a value of 1. The formula DistForce(n), or Dist(n ;1), will sample a new value from distribution number n each time the distribution is referenced in a tree calculation.

This also means that at each place in the tree where the distribution is referenced, a different sample value will be returned.

20.2.3 Defining distribution parameters non-numerically

Distribution parameters can be defined using variables or formulas, instead of fixed numeric values. This makes it easier to modify a distribution's parameters and it may also help someone viewing the model understand the significance of a particular distribution.



Using variables for distribution parameters

Clicking on the button to the right of a distribution parameter's text box will open an expression editor dialog. This dialog, like a variable definition window, makes it easier to set up a complex expression, including existing variables, functions, and even other distributions, to represent a distribution parameter.

Variables that are referenced in the parameter of a distribution must be defined with a default value for the tree, at the root node.



Distributions are "global" within the context of the model. Therefore, the root node variable definition will be used to define the distribution properties. You cannot define a variable differently in other portions of the model and expect the parameters of the distribution to change.

Microsimulation tracker variables (refer to the Individual-Level Simulation and Markov Models Chapter) can be used in the parameters of a distribution; however, the distribution must either be set to sample per Markov stage, or use DistForce() syntax, in order for a sample to be generated based on an updated value of the tracker.

Finally, it is possible to use one distribution ("X") in defining the parameters of another distribution ("Y"). TreeAge Pro requires that the dependent distribution have a higher numeric index than the input distribution – e.g., X=Dist(1), Y=Dist(2).

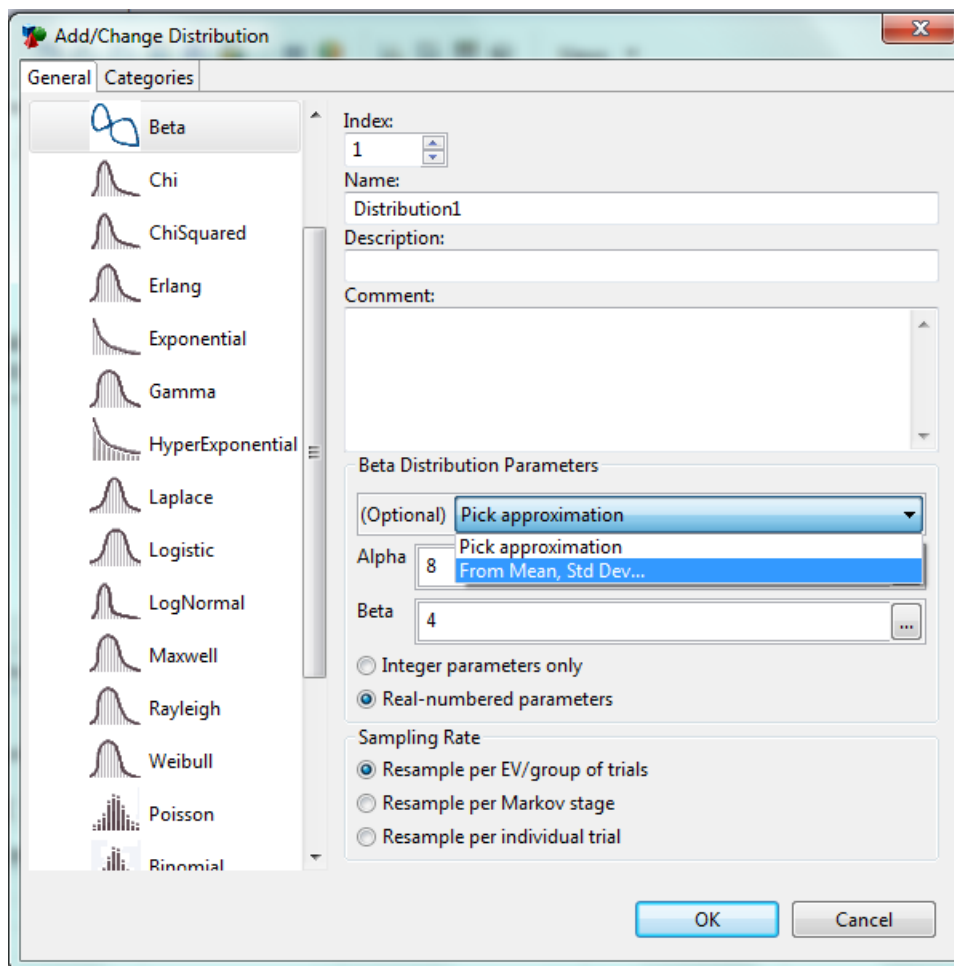
A distribution cannot reference itself recursively as a parameter.

20.2.4 Approximating distribution parameters from statistical information

Some of the built-in distribution types supported in TreeAge Pro use parameters that may be difficult to find. For example, the Log-normal distribution is parameterized using the “mean of the logs” and the “standard deviation or error of the logs”.

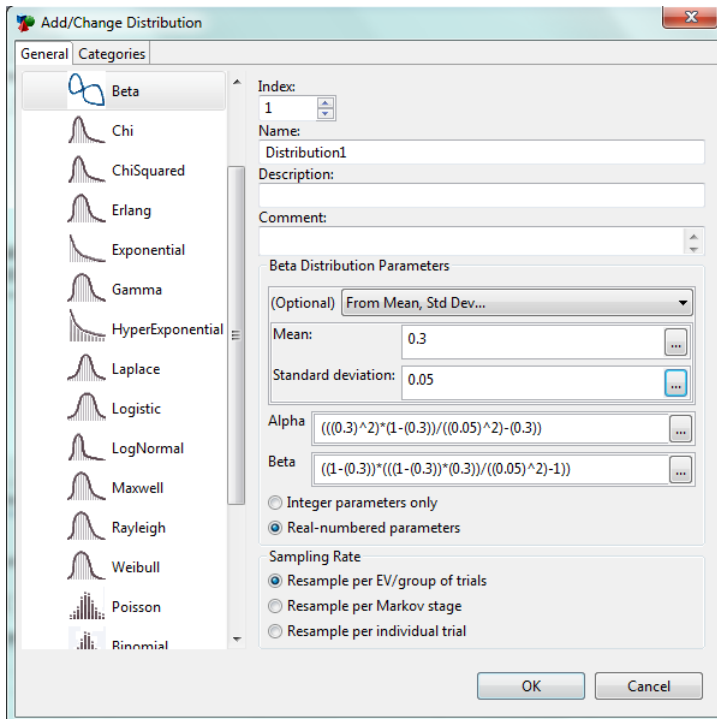
TreeAge can approximate some distributions’ parameters from more straightforward statistical summary values. Distributions that support parameter approximation will include a list box allowing you to approximate the distribution parameters. Supported distributions include the real-number form of the Beta, the Log-normal, and the Gamma distribution.

In the example below, beta distribution parameters are required.



Beta distribution - choose approximate

After choosing Pick approximation > From Mean, Std Dev..., the appropriate approximation inputs are presented for input entry. After entering the mean and standard deviation, the approximated distribution parameters are displayed.



Beta distribution - after entering approximation statistics

In a later section of this chapter, some of the details of these approximations are provided.

20.2.5 Correlated MultiNormal distributions

A model may have pairs or sets of distributions that should be positively or negatively correlated when sampled during a Monte Carlo simulation. The MultiNormal distribution type allows modelers to represent a multivariate normal distribution based on a set of correlations or variances/covariances.

A single MultiNormal distribution will sample multiple normally-distributed values. In reference to the MultiNormal distribution's individual, correlated sample values use the *Dist(index; variate)* function syntax.

The Special Features tutorial example tree "Multivariate Normal" illustrates the use of the MultiNormal distribution.

The first step in creating a MultiNormal distribution is to create the correlations. Set up a table with the correlations among the normal distributions. The following figure shows this data within the *Correlations* table.

Index	Value1	Value2	Value3	Value4
1	1	0	0	0
2	0.19	1	0	0
3	0.05	-0.05	1	0
4	-0.83	-0.66	-0.2	1

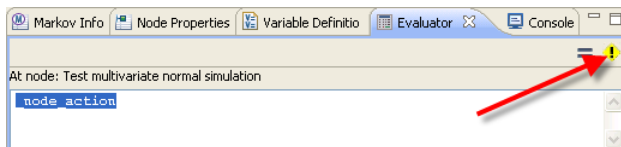
Correlations table

The correlation between distribution 1 and 2 is 0.19. The correlation between distribution 1 and 3 is 0.05, and so forth.

The correlation data must be manipulated via Cholesky Decomposition prior to using it in the MultiNormal Distribution. This is done via the Command function as illustrated in the `_node_action` variable definition.

```
_node_action = Command("TABLES"; "Correlations"; "CHOLESKYDECOMP"; "Cholesky")
```

This command preforms Cholesky Decomposition on the Correlations table and places the decomposed correlations in the *Cholesky* table. You can execute the `_node_action` by clicking on the exclamation point in the Evaluator view.



Evaluator view - execute `_node_action`

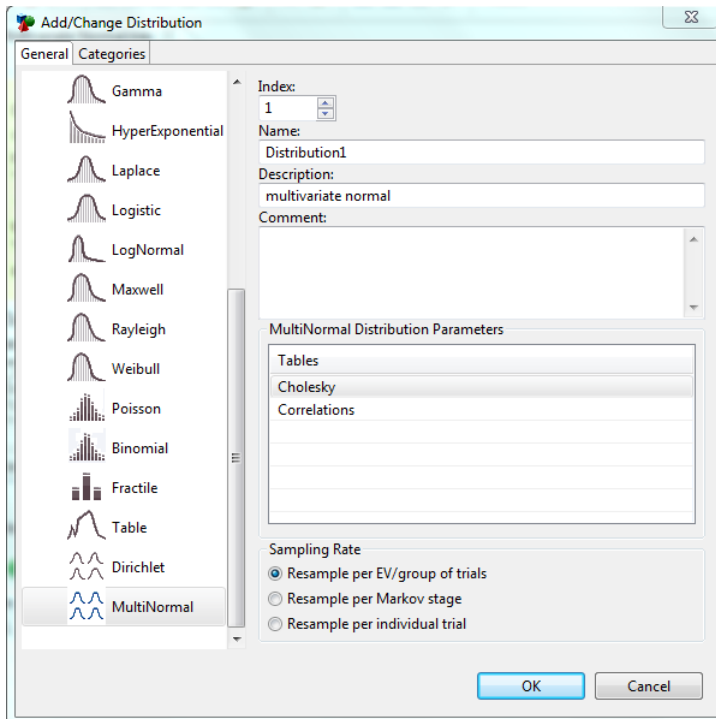
The following figure shows the decomposed correlations within the CholeskyCorrelations table.

Index	Value1	Value2	Value3	Value4
1	1	0	0	0
2	0.19	0.9817...	0	0
3	0.05	-0.060...	0.9969...	0
4	-0.83	-0.511...	-0.190...	0.1149...

CholeskyCorrelations table

Note that you can set the mean and standard deviation for each of the normal distributions by inserting two extra rows in the decomposed correlation table. This has no impact on the correlations. Since the example above has four normal distributions, you would place the mean and standard deviation for each in row 5 and 6, respectively.

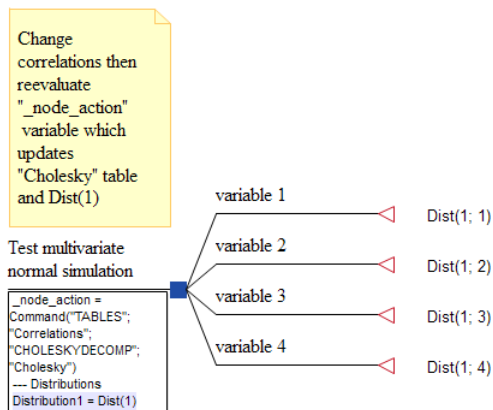
Once the correlations have been decomposed, they can be referenced in the MultiNormal distribution. The MultiNormalDistribution model contains such a distribution.



MultiNormal distribution

Note that the distribution type is MultiNormal and that the CholeskyCorrelations table is selected.

The distribution is now ready for use. The model itself samples each of the four normal distributions within the MultiNormal distribution.



MultiNormalDistribution model

Run a sampling simulation to review the correlated samples.

20.2.6 Correlation using Python user-defined functions

A different technique for correlation utilizes Python functions to access multivariate distributions in the “RandomArray” and other Numerical Python extensions. This requires a full installation of the free,

open source Python software. See the Python page at the TreeAge website, and the Numerical Python website:

<http://numpy.sourceforge.net/numdoc/HTML/numdoc.htm>

20.2.7 Creating user-defined Python distributions

User-defined Python distributions have not yet been implemented in TreeAge Pro.

20.3 Distribution Formulas

[Click here](#) to view the formulas TreeAge Pro uses to generate distributions.

20.4 Sampling from tables during Monte Carlo simulation

If it is not feasible or desirable to use one of TreeAge Pro's built-in distributions to represent the particular probability distribution you need, there are at least two ways to create custom sampling distributions. Both methods use a built-in distribution and a table.



Table distributions can be helpful in defining individual characteristics (i.e., gender, ethnicity) to trials during microsimulation based on the probability of each value within the cohort.

20.4.1 Creating a Table-type distribution

One way to sample values from a custom distribution is to create a new table in TreeAge Pro describing the distribution's discrete probability function (not the data set). This table can then be assigned to a Table distribution.

Each row of the table defines a distribution value (entered in the *index* column) and its probability (entered in the *value* of a table entry). The probabilities in the value column must total 1.0.

To create a Table distribution from an existing empirical data set:

- Use the Table Properties View to create a table and populate it with data that represents the custom distribution function. Refer to the Creating and Using Tables Chapter for details (see below).
- Within the Distribution Properties View, click the "add" toolbar icon to create a new distribution.
- In the Add/Change Distribution Dialog, select the distribution type "Table".
- Under "Distribution parameters", select the table from the list of the model's tables.
- Enter a name and description for the distribution.

- Click OK to save the distribution and close the Add/Change Distribution Dialog.

Index	Value1
1.0	0.25
2.0	0.5
3.0	0.25

Table for table distribution

Create Table Distribution - references table

To randomly sample from the Table distribution, simply reference the distribution in the standard way: use either the distribution's name or the `Dist(n)` function using the distribution's index within the model.



Sample values will only be drawn from exact table entry indexes, regardless of which lookup method you specify. TreeAge will not interpolate in a Table distribution.

The mean value of the Table distribution will be used as the distribution's expected value in non-Monte Carlo calculations.

Use the `Dist(index; 2)`, `Dist(index; 3)`, etc. distribution reference sequence to sample based on probabilities in the second, third, etc. value column.

20.4.2 Using a distribution to lookup values in a table

There are situations where distributions cannot be easily represented using a standard distribution type, or a regular Table distribution as described above.

For instance, perhaps you have a table of age-dependent probabilities or costs that you want to do a probabilistic sensitivity analysis on. Or, you might have a set of observed parameter values that you want to bootstrap from with each row's value given an equal probability (or even selected in order).

In these cases, you could populate the table and then use a separate distribution to sample row indexes and/or columns indexes.

For example, you could fill a table with observed data, numbered from 1 to N, and then pick values randomly from the table using a uniform distribution with a range equal to the range of table indexes. In this case, use the integer form of the Uniform distribution (to return only integers in the index range).

To sample from a table using a Uniform distribution:

- Enter/paste your data set into a TreeAge Pro table, using consecutive integer indexes in the index column and the data set's values in the value column.
- Within the Distribution Properties View, click the "add" toolbar icon to create a new distribution.
- In the Add/Change Distribution Dialog, select the distribution type "Uniform".
- For the low value, enter the lowest integer index from your table (i.e., 0 or 1). For the high value, enter the highest integer index from your table.
- Enter a name and description for the distribution.
- Click OK to save the distribution and close the Add/Change Distribution Dialog.

The actual reference in a tree formula should look something like the following:

TableX[Dist(1)]

where "TableX" is the name of the custom distribution table, and inside the square brackets is the reference to the Uniform distribution (assumed to have the index 1 in the example above). During a second-order simulation, the Uniform distribution will be resampled within its range, causing different rows from TableX to be drawn randomly with an essentially equal likelihood if done correctly.

It may also be possible to pick rows from the table *in order* during a simulation by using the keyword `_sample` (for sampling, use `_trial` for microsimulation) in place of the Uniform distribution. The `_sample` counter corresponds to the current iteration of the probabilistic sensitivity analysis simulation, incrementing by one at each resampling iteration.

To pick from a particular column in a multi-column table, simply add the appropriate column parameter to the table reference, such as:

TableX[Dist(1); 2]

To convert a time-dependent table of values for sampling, you might add additional columns that represent percentiles or bounds of each row. Then, you would use a distribution to sample a column index (i.e., between 2 and 3):

TableX[_stage; 2+Dist(1)]

21. Creating and Using Tables

This chapter provides instructions on creating tables of numeric values for use in custom sampling distributions and other tree calculations.

21.1 Creating and editing tables

TreeAge Pro can store indexed tables of numeric values that represent parameters in your model. Here are some basic facts about tables:

- Tables are typically created and stored with models in *.trex files. This is similar to how *.pkg files worked in previous versions.
- TreeAge Pro supports the use of global tables. However, the default location for global tables is no longer in a "Tables" folder in the TreeAge Pro application folder. Rather, each TreeAge document/project workspace includes a "Global Tables" project. This is the default location of example global tables (e.g., "tMort.tblx"), as well as *.tblx files automatically converted from a previous version's global *.tbl files.
- In TreeAge Pro, each tree has its own preferences related to the use of Global Tables. It is possible to create sub-folders in the Global Tables project specifically for one or a group of tree models, and to use tree preferences to point to the appropriate sub-folder. Or, a tree can be set to ignore the Global Tables project and only utilize tables contained within the model.
- Every table has an index column and a value column. A table can have additional value columns (up to 512) and from one row of values up to tens of thousands of rows.
- Each row's index value must be unique, but indexes are not required to be consecutive integers. TreeAge automatically numbers the value columns with integers (starts at 1).
- Tables can linearly interpolate values for missing indexes/rows, as well as columns.
- Only numeric values (no variable names or formulas) can be entered in a table.
- In Markov models, tables are often used to represent probabilities that vary over time (or other dimensions). Refer to the Building and Analyzing Cost-Effectiveness Models Chapter.
- A table can represent a parameter's empirical probability distribution to sample from during Monte Carlo simulation. (Refer to the Distribution Functions, Options and Types Chapter for details on sampling from tables and other distribution types.)
- Table Values can be loaded into a table in a variety of ways, including: paste tab-delimited text; edit via the Excel add-in; query an ODBC database; manually enter index-value combinations.

Facts about tables

21.1.1 Tables View

Tables are created and maintained via the Tables View.

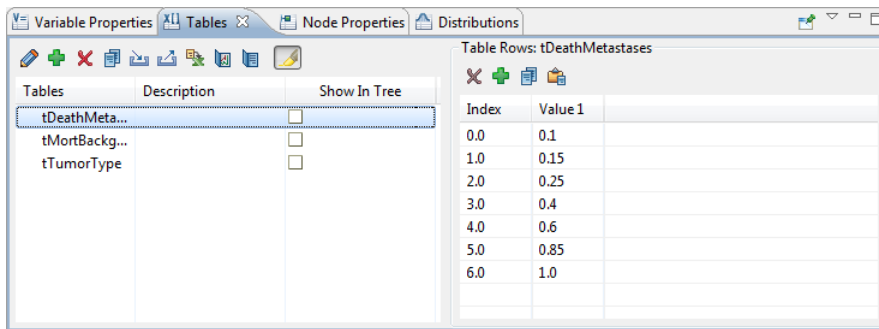
To open the Tables View:

- Choose Views > Tables from the toolbar.

Within the Tables View, there are two sets of data and associated controls

1. List of tables and table controls to the left.
2. List of table entries and table entry controls to the right.

When a table is selected in the list, the table entry data and controls apply to that selected table.



Tables View

The table property toolbar controls are described below from left to right.

- *Edit*: Edit the selected table from the list.
- *Add*: Create a new table.
- *Delete*: Deleted the selected table(s) from the list.
- *Duplicate*: Create a copy of the selected table and edit it as a new table.
- *Import*: Import table(s) from a global table file (*.tblx) into the model.
- *Export*: Export table(s) from the model to a global table file.
- *To Excel*: Copy the selected table and its data to Excel for editing.
- *Graph It*: Create a graph from the selected table's data.
- *Report*: Create the Tables Used Report.
- *Highlight*: Highlight the table within the model in the Tree Diagram Editor.

Table property toolbar controls

The table data toolbar controls are described below from left to right.

- *Delete*: Delete the selected table entries from the table.
- *Add*: Add a new table entry to the table.
- *Copy*: Copy table data with headers and rows to clipboard.
- *Paste*: Paste table data with headers and rows from clipboard to table. Existing data is replaced.

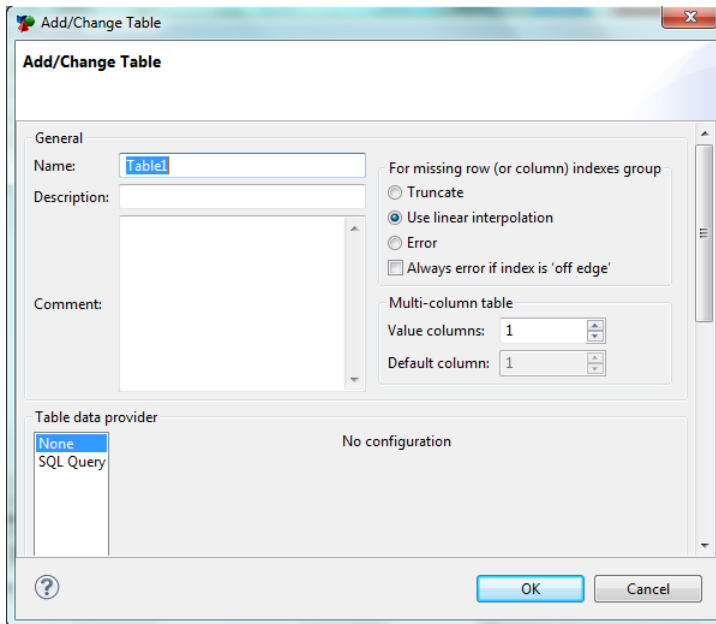
Table data toolbar controls

21.1.2 Creating a new table

Before entering values into a table, you must first create a table.

To create a new table:

- Open the Tables View.
- Click the "add" toolbar button. This will open the Add/Change Table Dialog.
- Enter the table name and properties into the dialog (see below). The specific property options will be described later in this chapter.
- Click OK to save the table and close the dialog.



Add/Change Table Dialog

After the table is created, it will appear in the tables list within the Tables View. You can then edit the properties of the new table.

To change the properties of a table:

- Select a table in the list.
- Click the "edit" button in the toolbar.
- Change the table properties in the Add/Change Table Dialog.

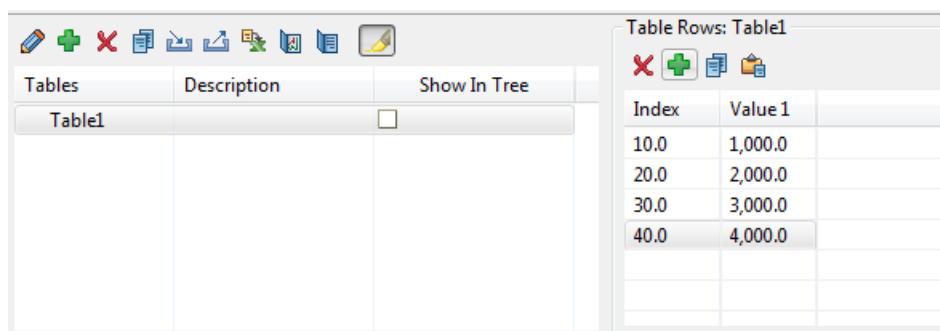
21.1.3 Entering values in a table

To populate a simple table, use the table entries toolbar and table data grid.

To enter table data:

- Select the table in the tables list.
- Click the "Add Row" button in the table entries toolbar. A new table row will appear in the table data grid.
- Change the values for the data row as needed.

- Repeat prior two steps to add more data rows.

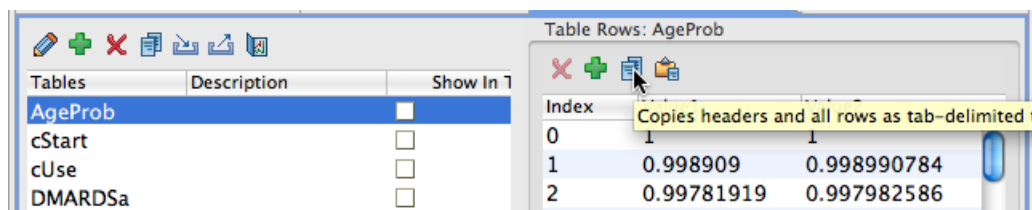


Tables View - with table and data

This method of entering data can be time consuming. There are other methods available for loading table data. Those methods are described in subsequent sections.

21.1.4 Loading/copying tab-delimited table data

To modify the values in an existing TreeAge Pro table, it may be more efficient to copy the table rows into a spreadsheet or text file for editing. Or, if you have a new, empty TreeAge Pro table, you can create the rows in a spreadsheet, and then copy and paste into the Table Rows list editor.



Tables View, Rows editor - Copy table data

To edit table data in Excel or a text editor:

- Select the table in the Tables view.
- Click the Copy button in the Table Rows editor, as shown above.
- Paste the rows into a spreadsheet or text document.
- Edit the data as tab-delimited text.
- Copy the table data, including headers, into the clipboard.
- Click the Paste button in the Table Rows editor.

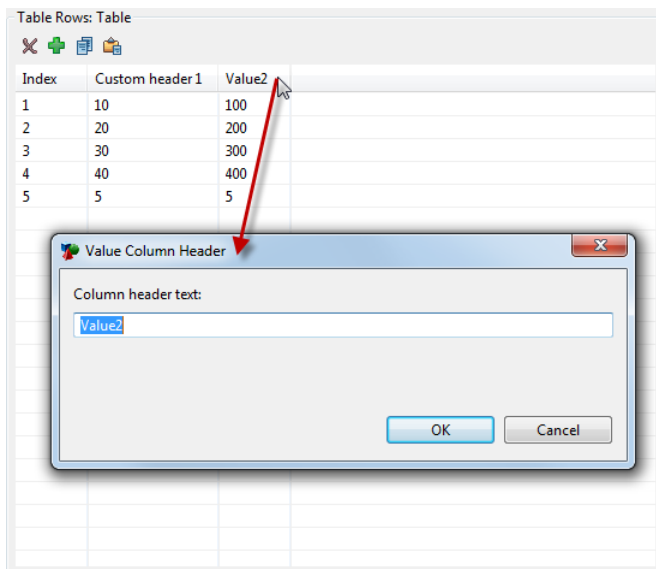
Take note of the following requirements:

- A column title row is not required. If one is included, however, the title of the index column must start with the word "Index". Value column titles have no restrictions.
- The selected range of cells should use either General, Text, or Number formatting (avoid accounting formats using parentheses for negatives numbers).
- The copied range should not include empty columns or rows.

- Instead of copying very large tables, it may be preferable to utilize a database to load data into a table, as described below.
- Note that the Paste Table command first removes existing rows from the table. However, the command is Undo-able.
- If the Paste command is not available, or does not work, see the bulleted list of suggestions above on proper cell formatting in the source document. If all else fails, try first pasting the spreadsheet data into a plain text file to remove formatting, and then copy the text file data into TreeAge.

21.1.5 Editing value column headers

You can enter custom column headers for value columns by clicking on the header. You can then enter the custom value column header into an input dialog.



Edit value column header

21.1.6 Editing table data in Excel

Users with the optional Excel Module can export table data to Excel, edit the data in Excel, then import the edited data back into TreeAge Pro. This technique is described in the Excel Module Chapter.

21.2 Referencing tables in formulas

There are two basic ways to use tables:

1. In order to look up a value in the table from a payoff, probability, or other calculation, you will need to use the proper syntax (below).

2. If a table is used as a sampling distribution, simply select its name from a list of tables when setting up the distribution (described later in this chapter).

21.2.1 Table lookup syntax

In a table composed of index-value pairs (i.e., a single value column), a value can be retrieved with the following syntax:

TableName[index]

The table's variable-type name is followed immediately by square brackets containing an index used to pick a table row. The index expression can be a number, a variable, or even another table reference.

If a table has more than one value column, specify the number of the value column to pick from (after the row index argument):

TableName[index; value column #]

The value column # expression can also be a number, variable, etc. However, this expression normally resolves to an integer value \leq the number of columns in the table. Note that when looking at the table, the first value column is actually the second column in the table because the first column contains the index.

For tables with more than one value column, table lookups without a value column # provided will use the default value column defined in the table properties.



Reverse table lookups

Special syntax for the `Command()` function enables reverse table lookups. This is designed to facilitate, for example, using a `uniform[0,1]` random number to "sample" age-at-event table from an age-indexed table whose value columns describe one or more inverse "survival" functions, or "cdfs". The syntax is:

`Command("TABLES"; "t_events"; "ReverseLookup"; prob; value_column)`

Other methods are: "ReverseLookupInterpolate", "ReverseLookupTruncate", and "ReverseLookupCeiling". Refer to the Tools and Functions for Complex Trees Chapter for additional information.

21.2.2 Table lookup methods

During tree calculations, if a formula references a table using an index value that matches a row index in the table exactly, `TreeAge` simply returns the value from the appropriate column in that row. However, when a reference is made to a non-existent row/index, the table's missing row lookup method determines what is returned.

The missing row/column lookup methods are:

- *Truncate*: If rows with lower indexes exist, returns the value from the row with the highest index that is less than or equal to the requested index; otherwise, returns the value from the first row (i.e., the lowest index value).
- *Use linear interpolation*: Returns a value calculated by linear interpolation between existing indexes. This is the default lookup method.
- *Error*: Report an error if an index value is provided that is not in the table.

Table lookup methods

Each table also has an “Index off edge is error” option. If this option is left unselected, a table that uses truncation or interpolation will allow references to indexes above or below the table’s actual range. In these cases, the table will return the value associated with the closest existing row or column. If this option is checked, on the other hand, an error will be reported.

TreeAge Pro supports interpolation of both rows and columns. The table below is used to show how TreeAge Pro handles table lookups with interpolation.

Index	Value1	Value2
10.0	1,000.0	10,000.0
20.0	2,000.0	20,000.0
30.0	3,000.0	30,000.0
40.0	4,000.0	40,000.0

Table interpolation example

Interpolation starts with the value for the index below the provided value. Then it adds the difference between the next higher value and the lower value based on the proportion of the way from the lower index to the higher index.

$$\text{TestTable}[14;1] = \text{TestTable}[10] + ((14-10)/(20-10)) * (\text{TestTable}[20] - \text{TestTable}[10])$$

$$\text{TestTable}[14;1] = 1,000 + (4/10) * (1000) = 1,400$$

$$\text{TestTable}[30; 1.3] = 3,000 + ((1.3-1)/(2-1)) * (30,000-3,000) = 11,100$$

TreeAge Pro supports simultaneous interpolation of both rows and columns. First interpolation is done on the row values and then those row interpolated values are used for the column interpolation.

$$\text{TestTable}[16;1] = 1,600$$

$$\text{TestTable}[16;2] = 16,000$$

$$\text{TestTable}[16;1.4] = 1,600 + ((1.4-1)/(2-1)) * (16,000-1,600) = 1,600 + (.4/1) * 14,400 = 7,360$$

21.3 Model tables and global table files

In TreeAge Pro 2009 and earlier versions, tables were saved by default as independent *.tbl files in a "Tables" directory found in the application folder. These global tables could be accessed by multiple models. It was also possible to then save a tree as a *.pkg file which contained copies of the tables. The default behavior has changed a little to be more flexible.

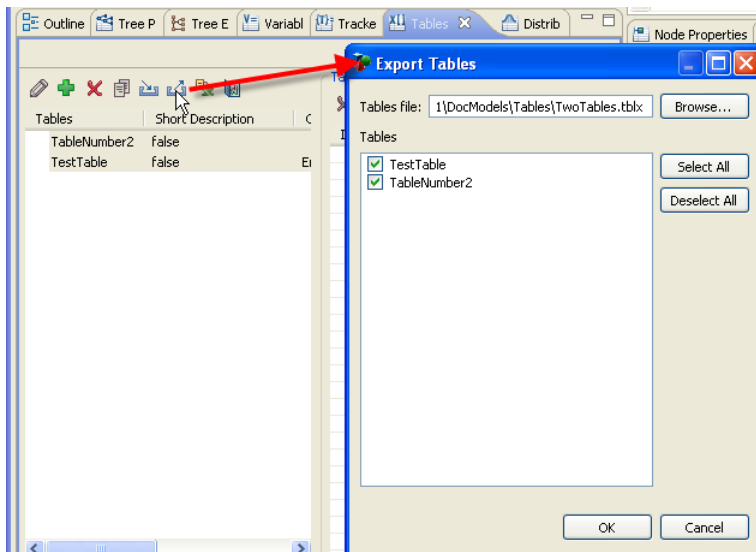
Tables are now normally created within a model, using the Tables view as described above, and stored within the model document (*.trex files). It is still possible to create and use global tables, which are now stored in a Global Tables project in your TreeAge workspace (instead of a program files sub-directory). When a new TreeAge project workspace is created (for example, when you run a newer TreeAge Pro for the first time), old *.tbl files found in a previous TreeAge Pro installation are copied into *.tblx files in a new Global Tables project. Tables can be easily imported and exported between and among *.trex models and Global Table files (*.tblx), which can exist outside the context of any model. Additionally, each *.tblx file can hold multiple tables.

The next sections demonstrate how to export tables to and import tables from a global table file.

21.3.1 Exporting tables from a model to a global table file

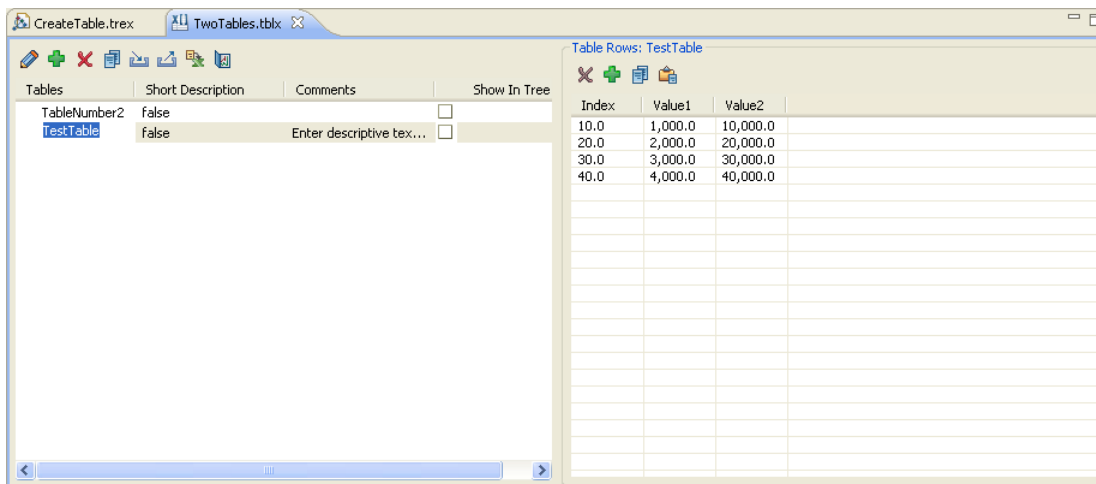
To export tables:

- Open a model with tables.
- Choose Views > Tables from the toolbar.
- Select the tables you wish to export.
- Click the "Export Global Tables" icon from the Table Properties toolbar.
- In the Export Tables dialog, specify the tables file and select the tables you wish to export.



Export tables to global tables file

The global tables file can then be opened as you would open a tree document. When opened, the global tables file will appear in the Tree Diagram Editor. However, its appearance will be very similar to the Tables View for the original model.

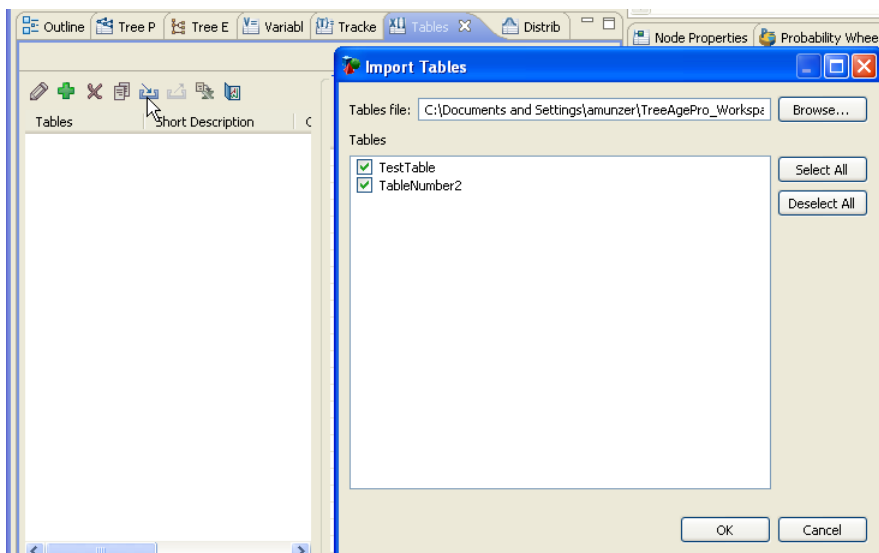


Global tables file

21.3.2 Importing tables from a global table file to a model

To import tables:

- Open a model.
- Choose Views > Tables from the toolbar.
- Click the "Import Global Tables" icon in the Table Properties toolbar.
- In the Import Tables Dialog, select the global tables file.
- Select the specific tables from the global tables file that you want to import.



Import tables from global tables file

The tables will then be included in the model and can be referenced in expressions.

21.4 Importing tables from older versions of TreeAge Pro (2009 and earlier)

You may have TreeAge Pro tables that were created in TreeAge Pro 2009 or an earlier version. Rather than re-entering the table properties and data, the table data can be imported into the current version of TreeAge Pro.

If you look in the Project view, there should be a Global Tables project. This may contain *.tblx files created by automatically importing older *.tbl files found in a previous version's "Tables" sub-directory. (Each time a new workspace is created, #2 below is automatically performed.)

A *.trex model can utilize the tables contained within a *.tblx Global Table for analysis purposes. These tables can be edited by double-clicking on the *.tblx file to open a tables editor.

There are a number of other options for importing table data into a tree model or a Global Tables project or folder, for example:

1. Import directly into *.trex model from a v200x *.pkg packaged tree, or from a previous version's "Tables" sub-directory.
2. Create a single *.tblx file containing all global tables from a v200x *.pkg, or the v200x "Tables" sub-directory.
3. Create individual *.tblx files for each global tables from a v200x *.pkg, or the v200x "Tables" sub-directory

In TreeAge Pro 2009 and earlier, by default, tables were written to global table files (*.tbl) stored in a specific folder on your computer. You also had the option of saving your model as a package file (*.pkg), which would include the tables used by the model within the model document itself.

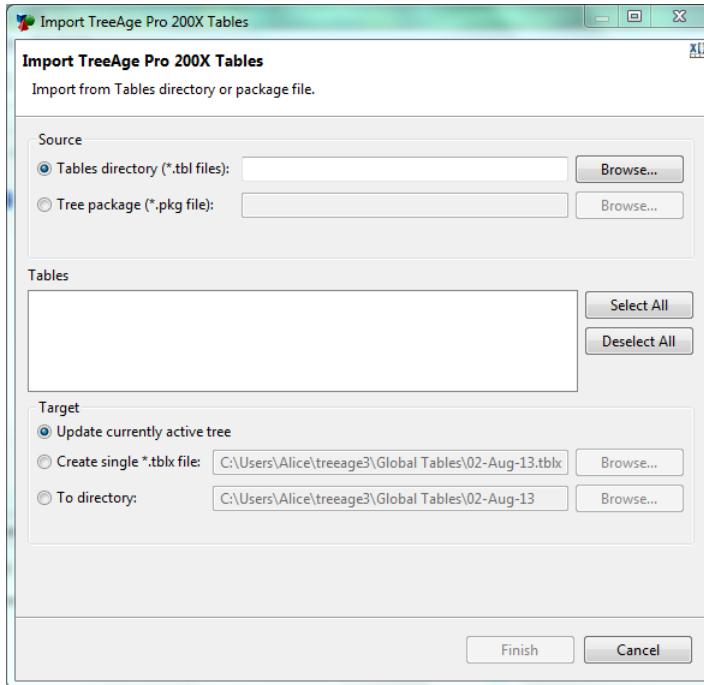
If you open an older package file in the latest version of TreeAge Pro, it will convert both the model and its tables into a current TreeAge Pro model file (*.trex). No specific action is required for the tables because the tables that were included with the original 2009 package file are also included with the current model file.

However, when you open a v200x tree file (*.tre), any referenced tables will not be converted with the model because the tree file does not contain table data. However, the required *.tbl files may already have been imported as *.tblx files in the Global Tables project in your workspace. If this is the case, analyses of the imported tree may work immediately without further action. If required *.tbl files were not accessible (e.g., previous version was on a different computer), you have two options: save the v200x tree file as a package file before conversion, or manually import the global *.tbl table files required by the tree file.

You can import all the 2009 table files (*.tbl) at once from the 2009 global table directory or you can import the tables contained within a 2009 package file (*.pkg). In either case, the tables can be imported either directly into the active tree document, or into a global table file.

To import 2009 tables:

- Click the "Import Tables" button in the Tables view toolbar, and then click the button labeled "Import v200x Tables"; or
- Choose File > Import/Export > Tables Import...
- The Import TreeAge Pro 200x Tables dialog will open.



Import TreeAge 200x Table Files Dialog

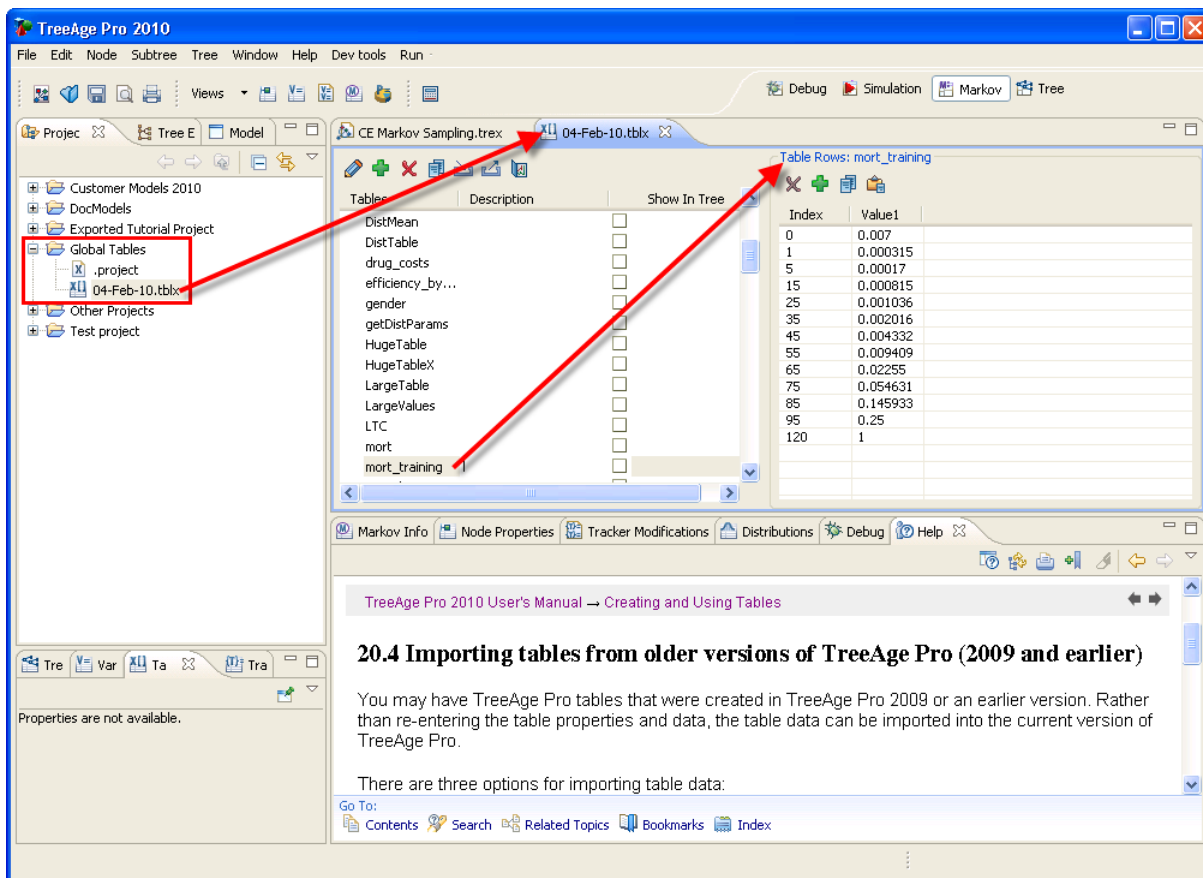
This dialog provides options for selecting the location of the table's source, the list of tables, and the target to receive the tables.

Let's say you want to import some or all of the global tables from your previous TreeAge Pro 2009 installation into the currently open tree document, which perhaps was imported earlier from a v200x *.TRE file. In the dialog you would...

1. Browse to a Tables folder from a previous TreeAge Pro v200x installation. (The default path, if it exists on the computer, will be entered for you.)
2. Select the desired tables to import.
3. Use the default, selected target, which will import into the currently active tree.

After you click Finish, the open tree document's list of tables will be updated to include the imported tables.

Or, instead, you can choose to import the tables into a new or existing Global Tables file for modification and eventual re-import into a Tree Document. After you click the Finish button, the import is executed; if necessary, a new Global Tables project will be created in your TreeAge Pro Workspace. Within the Global Tables project, the *.TBLX files will be updated/created.



Global Tables project with imported tables

To make changes to the imported tables, open the Tables document file (e.g., "04-Feb-10.TBLX") in the Global Tables project. When tables contained in the Tables document are needed for a particular tree, simply go to the tree document's Tables view and import tables from the tables document.

Variations on the import described above include:

- Import 2009 tables from a 2009 package file (*.PKG) rather than the v200x Tables folder.
- Import 2009 tables to an existing global tables document (*.TBLX) rather than a new tables document.
- Import 2009 tables to a folder rather than a single tables document; each table is contained in a separate Global Tables document within the folder.

21.5 The All Tables Report

The All Tables Report displays all the tables that exist and/or are referenced in the tree.

Name	Description	Unused	Missing	Comment
MyTable2	t2-desc	UNUSED		t2-comment
MyTable1	t1-desc			t1-comment
MyTable3			MISSING	

All Tables Report

The "Unused" column indicates that a table exists in the tree but is never referenced. The "Missing" column indicates that a table is referenced in the tree, but does not exist.

21.6 Managing tables using the Excel module

Users with the Excel/COM module can utilize the TreeAge add-in menu in Excel to update TreeAge with changes made to tables stored in a spreadsheet. Refer to the Graphing, Reporting and Modeling Using Excel Chapter.

It is also possible to use the TreeAgeProLib to script changes to tables. Refer to the Using the TreeAge Pro Object Interface Chapter.

21.7 Linking a table to an ODBC data source

TreeAge Pro includes the option to link a table to an existing database or other data source that has an ODBC driver. Linking a table to a database is an easy way to make table updates automatic.

TreeAge Pro tables can use ODBC to access most commercial and open-source database formats, including: MySQL, SQL Server, Access, Firebird, as well as Excel spreadsheets, text-format files, and XML files. A standard SQL SELECT query is used to select columns and rows from a table in the data source, which then replace the contents of the target TreeAge Pro table.

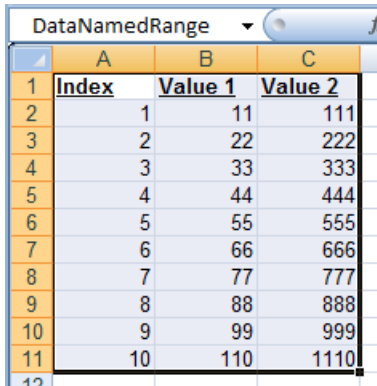
21.7.1 Using ODBC data sources in Windows

Before you can attach a TreeAge Pro table to an ODBC data source, you must first create the ODBC connection to the data source. Microsoft has published instructions for on ODBC data sources on their website (Windows XP, Windows 7).



Creating ODBC data sources has become more complex with Office 2007 and Windows 7. Please refer to Microsoft documentation regarding creating ODBC data sources. Note that this example uses the 32-bit ODBC driver setup on a Windows 7 computer since no Office ODBC drivers were listed under the 64-bit Data Sources entry in the Control Panel.

In this document, we will create an ODBC data source in Windows XP that connects to the following Excel worksheet saved on the computer.



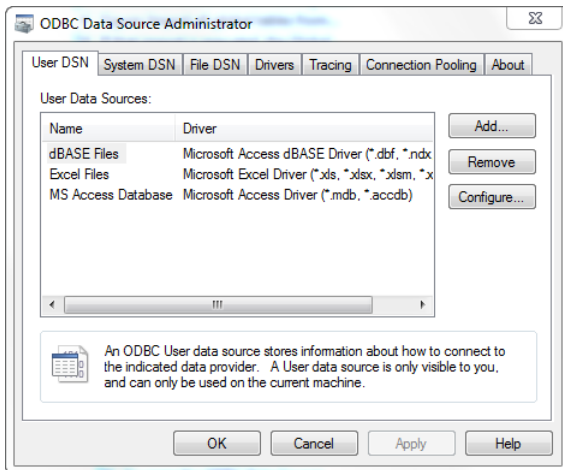
	A	B	C
1	Index	Value 1	Value 2
2	1	11	111
3	2	22	222
4	3	33	333
5	4	44	444
6	5	55	555
7	6	66	666
8	7	77	777
9	8	88	888
10	9	99	999
11	10	110	1110

Excel data source

Note that the data is contained within the named range *DataNamedRange*.

To open the ODBC Data Source Administrator in Windows 7 via the 32-bit drivers:

- Open the 32-bit ODBC Data Source Administrator by executing the program C:\Windows\SysWOW64\odbcad32.exe.



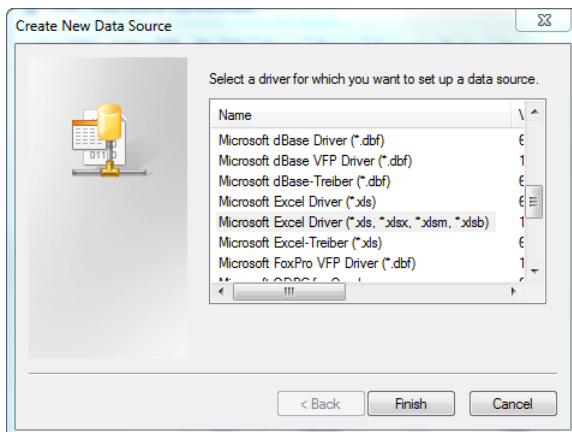
ODBC Data Source Administrator Dialog

System data source names (DSN) are available to all users of the computer, so we will create a System DSN.

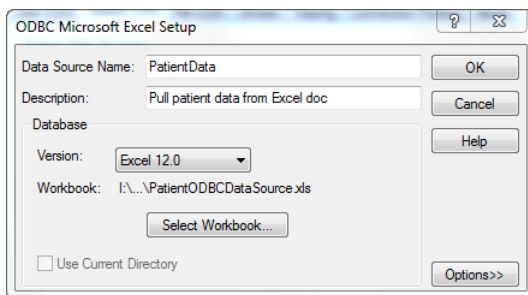
To create a System DSN:

- Select the System DSN tab.
- Click the Add button.
- Select the Microsoft Excel driver from the list presented in the Create new Data Source Dialog (see below).

- Enter a data source name and select the workbook in the ODBC Microsoft Excel Setup Dialog (see below).

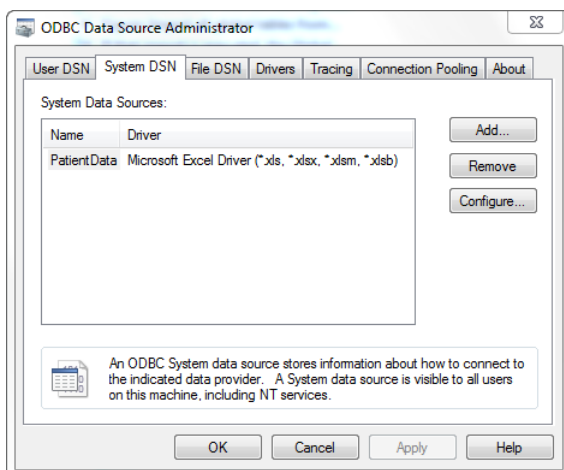


Select ODBC driver



Select workbook

After creating the System DSN, it will appear in the ODBC Data Source Administrator Dialog.



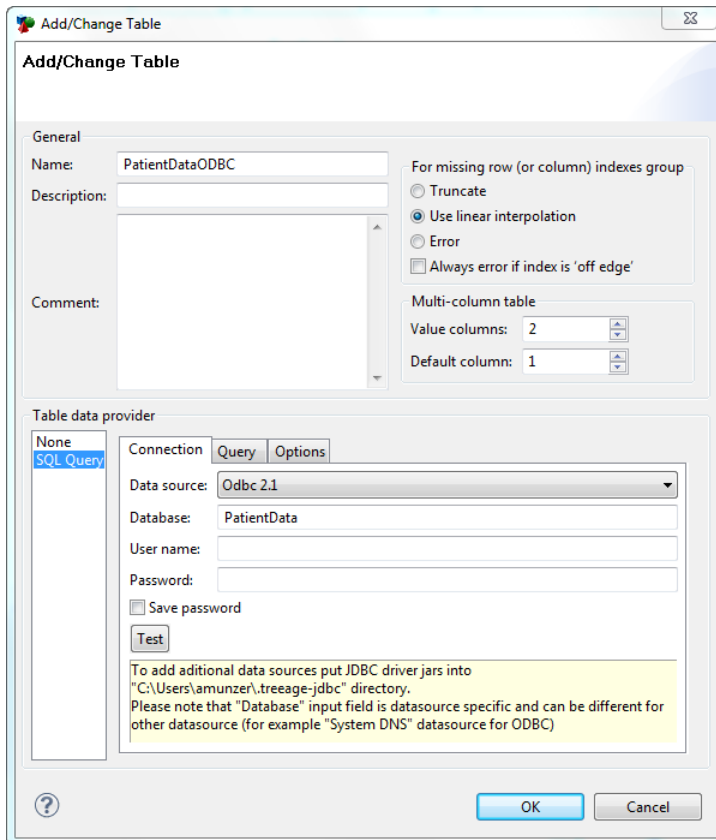
ODBC Data Source Administrator - note new source PatientData

Now that the ODBC data source is ready, we can connect it to a TreeAge Pro table.

To connect a TreeAge Pro table to an ODBC data source:

- Create a table via the Tables View.

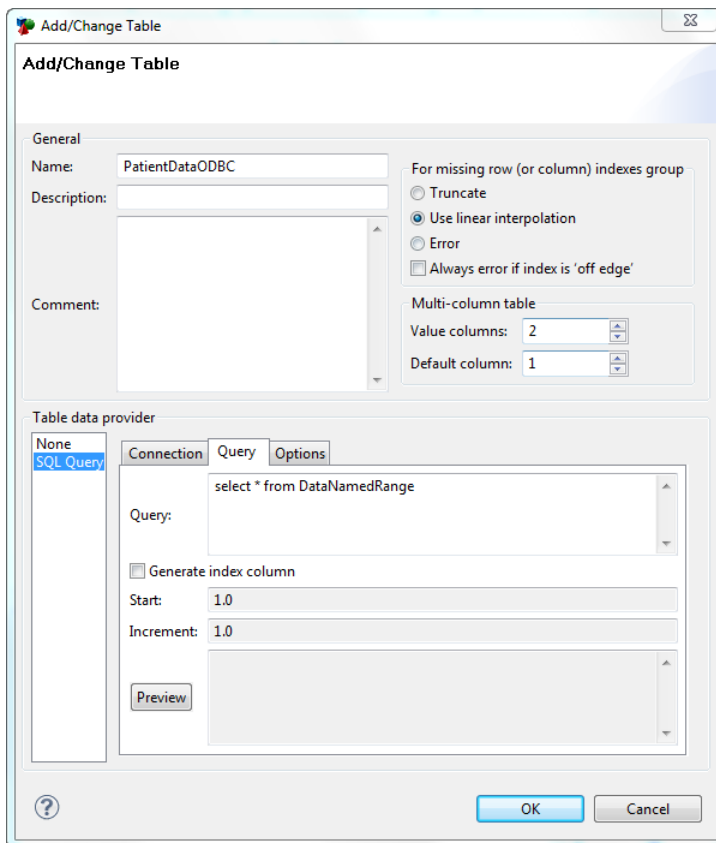
- In the Add/Change Table dialog, click on the SQL Query option in the Table data provider area.
- For Data source, enter the value "ODBC 2.1".
- For Database, enter the name of the data source from the ODBC setup.
- Click Test to verify that the connection can be established.



Setup table with ODBC data source - 1

To connect a TreeAge Pro table to an ODBC data source (continued):

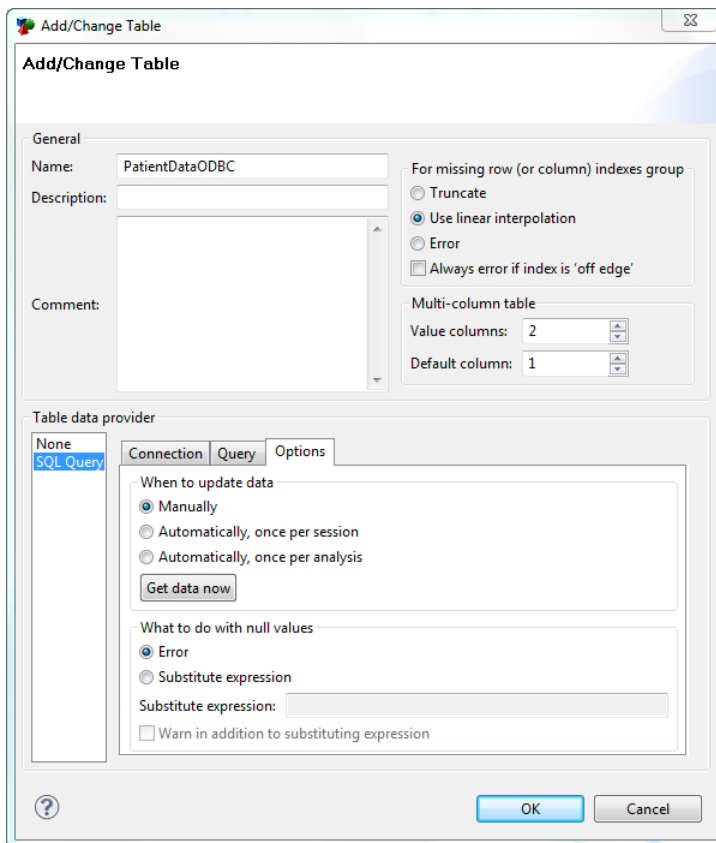
- Click on the Query tab.
- Enter the query expression into the appropriate field. This example refers to the name data range in the Excel document. You can also refer to a worksheet within the document.
- Use the other options as needed.



Setup table with ODBC data source - 1

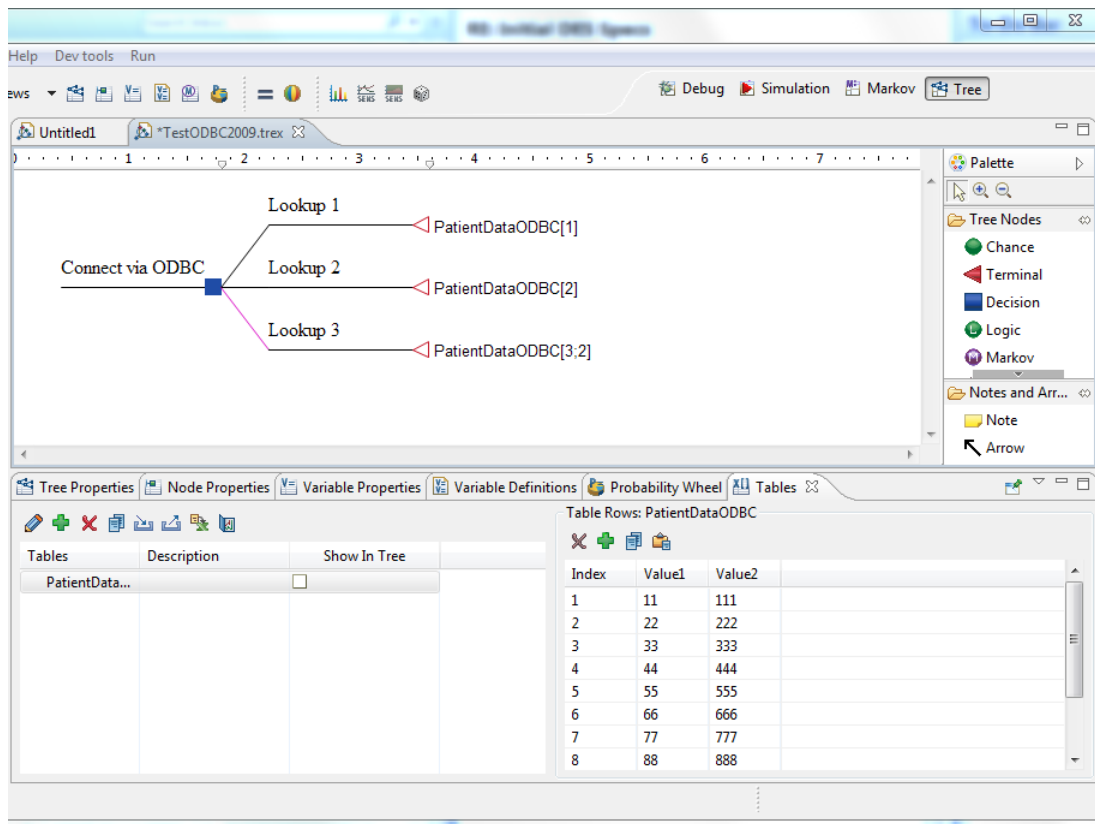
To connect a TreeAge Pro table to an ODBC data source (continued):

- Click on the Options tab.
- Enter options for when to load data and what to do with missing data from the data source.



Setup table with ODBC data source - 1

The table is now available for use in your model.



Model with ODBC table

22. Stored Analysis Abstracts and Sequences

TreeAge Pro allows you to store the settings used to perform many types of analyses. This chapter provides instructions on saving and using stored analyses. It also describes how to create simple stored analysis sequences that, for example, runs a series of simulations or sensitivity analyses.

For more complicated tasks, such as batching lengthy analyses, automating export of analyses to spreadsheets, or retrieving analysis parameters from a database or spreadsheet, you can use TreeAge Pro's Object Interface.

22.1 Using stored analyses

It is possible to save and reuse the parameters for most of the analyses available in the Analysis menu. This includes Monte Carlo simulation and all forms of sensitivity analysis. Analyses that cannot be stored include Graph Risk Preference Function, Show Optimal Path, Verify Probabilities, and Roll Back.

22.1.1 Create a stored analysis

To create a stored analysis, you must first run a regular analysis. TreeAge Pro can only save the last analysis, so you must save the stored analysis before running a second analysis on the model.



Only certain analysis types can be stored - Monte Carlo simulation, sensitivity analysis and tornado diagrams.

For this example, we will save a stored analysis for a Monte Carlo simulation with trials on the Three Vars Get Started tutorial example model.

To store the parameters of an analysis:

- Open the Three Vars example model.
- Select the root node.
- Choose Analysis > Monte Carlo simulation > Microsimulation (Trials) and start the simulation.
- Either cancel the analysis or allow it to complete.
- Select the tree in the Tree Diagram Editor.
- Choose Analysis > Stored Analyses from the menu. This opens the Tree Preferences category Stored Analyses.
- Click the Save Analysis button.
- Change the default name to something meaningful (see warning below) and click OK.
- Save the model.

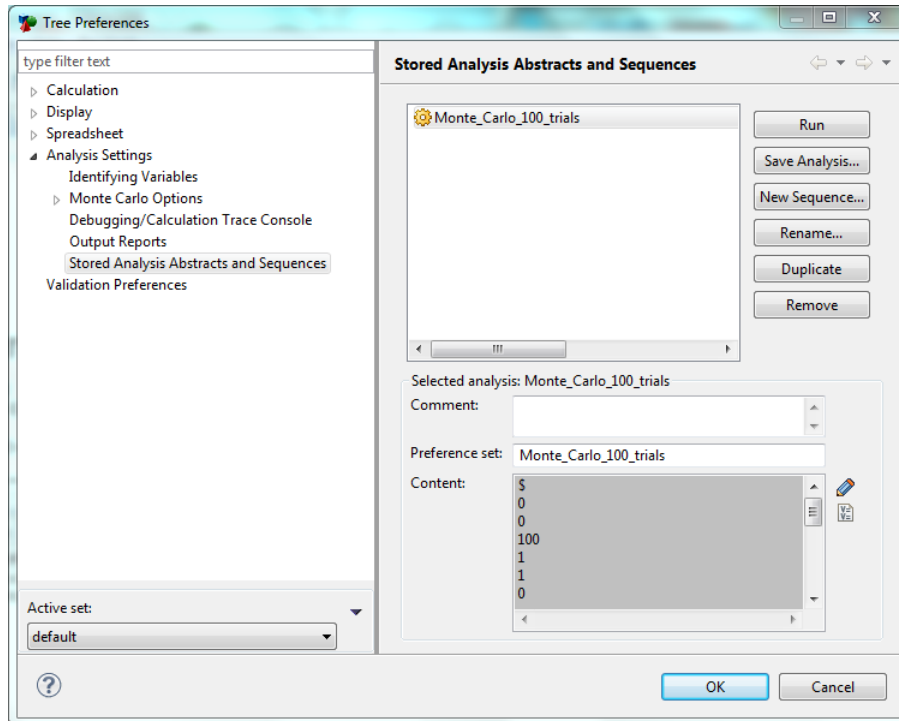


Use only letters, numbers and spaces in your stored analysis name if you want to use the analysis in a stored analysis Sequence. Sequences cannot handle punctuation.



If you are storing a Monte Carlo simulation to reuse, you do not have to let the simulation run to completion.

The instructions that TreeAge Pro needs in order to perform the exact analysis are stored in the Tree Preferences (the results of the analysis are not).



Tree Preferences - Stored Analyses

The top of the dialog shows a list of stored analyses and sequences. The bottom of the dialog shows the parameters used by the selected stored analysis.

22.1.2 Run a stored analysis

Once an analysis is stored, it is saved with the tree, and can be run as long as no significant structural changes are made where the analysis was originally run.

To run a stored analysis:

- Open the tree in which you have stored an analysis.
- Choose Analysis > Stored Analyses from the menu. This opens the Tree Preferences category Stored Analyses.
- Select a stored analysis from the list.
- Click Run.

Many analyses are dependent on the structure and variable definitions of your tree. For instance, the location (relative to the root node) of the node at which you originally performed a sensitivity analysis

must be unchanged from when you stored the analysis, and all analyzed variable definitions must be present in the same locations. If TreeAge Pro is unable to reconcile the different structures, it will not run the analysis.

22.1.3 Edit/maintain stored analyses

The Tree Preferences category Stored Analyses allows you to edit, rename, delete or copy a stored analysis.

To edit a stored analysis:

- Select an existing stored analysis.
- Edit the comment if desired.
- Click the pencil icon to edit the stored analysis parameters if desired.



Be very careful when editing stored analysis parameters since they are not labeled. In the example above, you could safely change the 100 to a different number of trials. In the case of a sensitivity analysis, you could edit a variable and/or its range.

You can also rename or duplicate an existing stored analysis using the appropriate buttons to the right of the stored analysis list.

22.2 Sequencing stored analyses

In TreeAge Pro, sequences of stored analyses can be defined and stored, like a batched analysis, for future use. Analysis sequences can greatly simplify tasks like batch processing a series of lengthy analyses, such as Monte Carlo simulations. Sequences can also be used, if you frequently do manual kinds of sensitivity analyses, by repeating any of TreeAge Pro's built-in analyses over a range of variable values.

22.2.1 Creating an analysis sequence

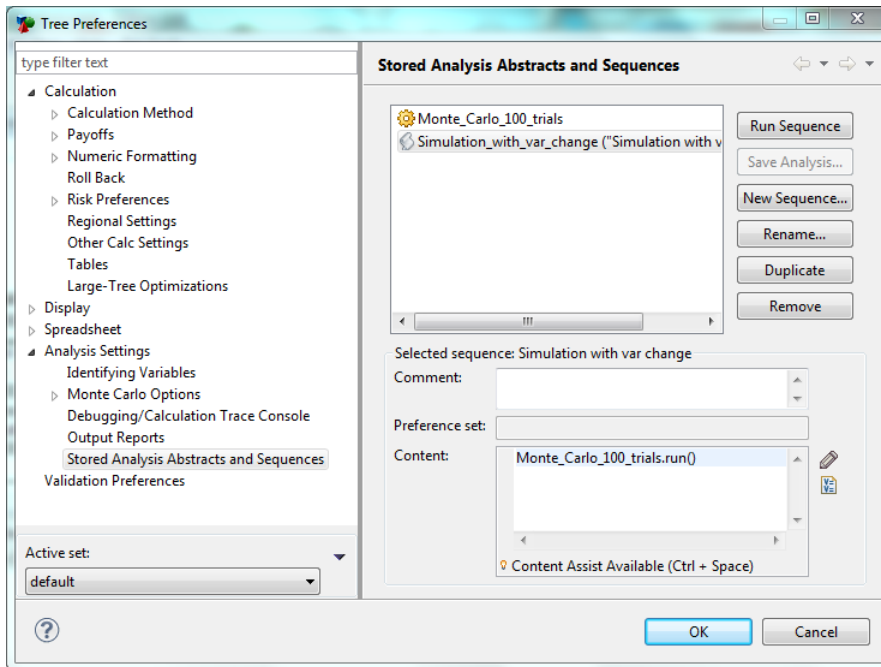
This section provides a simple illustration of how the sequencing feature can be used to automate running a series of separate Monte Carlo simulations, each using a different value of a variable. We will assume that you already created the original Microsimulation stored analysis described earlier in this chapter.

To create a sequence of stored analyses:

- Choose Analysis > Stored Analyses from the menu.
- Click the New Sequence button.
- Enter a name for the new sequence (Simulations with var change).
- Select the new sequence in the stored analysis list.
- Click in the first line of the Content input list.

- Press **CONTROL+SPACEBAR** to use autofill to bring up a list of existing stored analyses.
- Select the Monte Carlo simulation stored analysis.

At this point, you will have a sequence that simply runs a single stored analysis (see below).



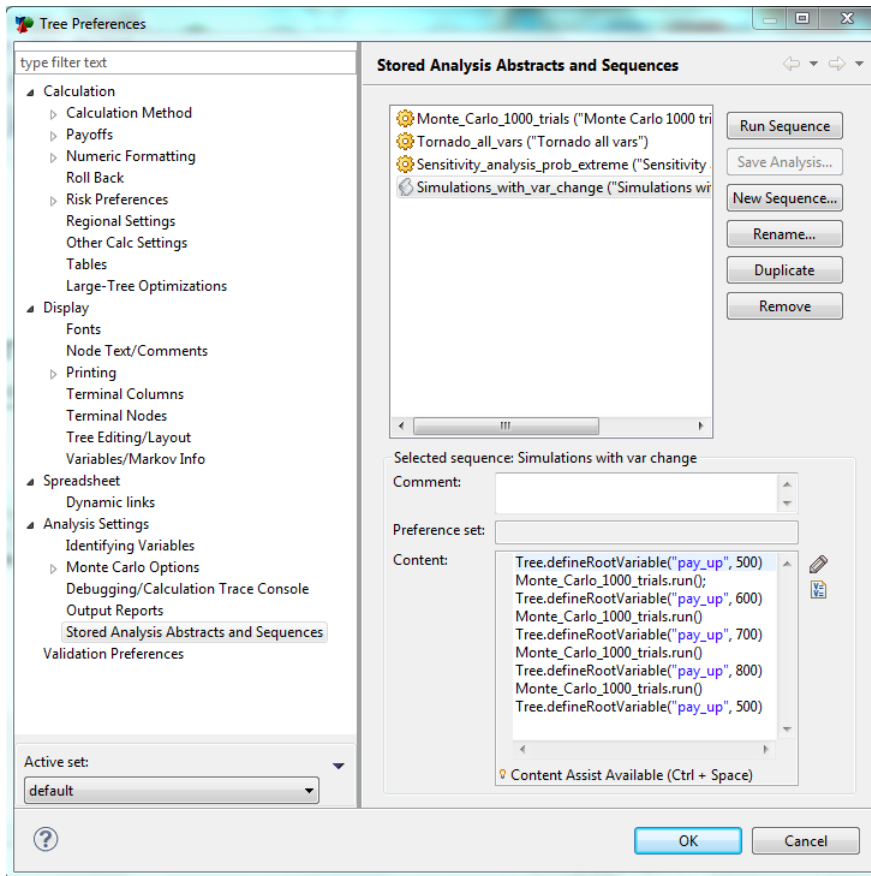
Stored analysis sequence

In addition to stored analyses, you can also make minor modifications to the tree via sequence commands. The most common modification is to change the value of a variable at the root node. This is achieved through the following command:

Tree.defineRootVariable(variable, definition)

Autofill can help create the sequence command.

The Special Features tutorial example model, Three Vars with Stored Analyses, contains three stored analyses and a sequence which runs four microsimulations, each with a different value for the pay_up variable.



Stored analysis sequence

22.2.2 Run/maintain analysis sequences

You can run, rename and delete a stored analysis sequence using the same buttons as for individual stored analyses.

Stored analysis sequences can be edited by selecting the sequence, then editing the Content field.

23. Linking with Excel and Other Applications

This chapter covers dynamic linking, a powerful tool for integrating a decision tree and an Excel spreadsheet.

23.1 Dynamic linking with Excel via BiLinks

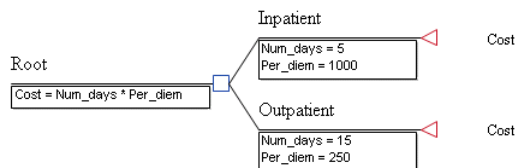
If you have developed a complex set of cost or utility calculations in a spreadsheet, rather than trying to recreate the formula in a decision tree, you can create dynamic links between the tree and the spreadsheet. With dynamic links, TreeAge Pro inputs a set of variables into corresponding spreadsheet cells, recalculates the spreadsheet, and retrieves results from the output cells.



Note that the flow of data is from TreeAge Pro to Excel back to TreeAge Pro, not vice versa.

Dynamic links work during any type of analysis (e.g. roll back, sensitivity analysis, and Monte Carlo simulation).

To illustrate the basic workings of dynamic links, consider the Bilinks tutorial example tree BiLink-before.trex pictured below.



BiLink-beore.trex Tree

The Cost variable is simply defined with the formula “Cost = Num_days * Per_diem” at the root node. However, you might have a highly complex set of calculations to generate the value of Cost; these calculations might be more easily done in Excel. This technique is illustrated by placing this simple multiplication formula in the Bilinks tutorial examples Excel worksheet BiLink-worksheet.xls. See below.

	A	B
1		
2	Days of treatment	15
3	Cost per day	250
4		
5	Total cost	=B2*B3

BiLink-worksheet.xls Worksheet

The variables that are inputs into the payoff calculation, Num_days and Per_diem, are given different definitions at each terminal node. These definitions will be passed to the spreadsheet, which will then recalculate the product and pass it back to the Cost payoff variable.

23.1.1 Creating a BiLinks report

A BiLinks report is useful if you have many dynamic links in your model. The BiLinks report details inbound BiLinks from Excel and outbound BiLinks to Excel.

The report provides the following information about inbound BiLinks from Excel:

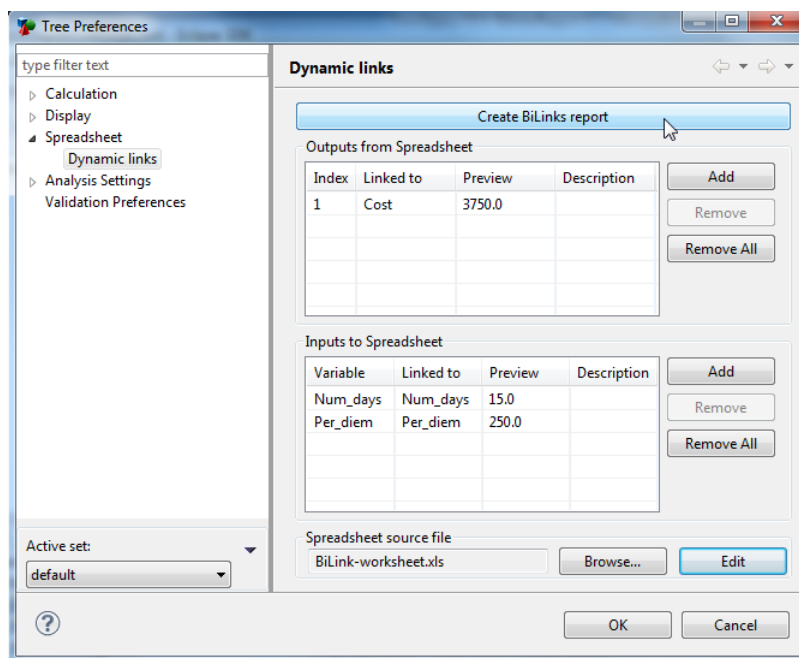
- BiLink Index
- Linked Excel cell name
- Value from Excel
- TreeAge variable, if the variable is defined at the root node
- Description, if a description is provided from the BiLink entry.

The report provides the following information about the outbound BiLinks to Excel:

- Linked TreeAge Pro variable
- Linked Excel cell name
- Value to Excel
- Description, if a description is provide from the BiLink entry.

Using the BiLinks tutorial example, BiLinks-after.trex, use the following steps to generate a BiLinks report:

- Open Tree Preferences > Spreadsheet > Dynamic Links, as indicated in the Figure below.
- Select the button "Create BiLinks report" to generate the report.
- To change any of the outputs to and inputs from the spreadsheet, see the next section Calculating payoffs using Dynamic links.



Tree Preferences - Generating the BiLinks report

Inbound Bilinks from Excel				
Index	Excel Cell	Excel Value	TreeAge Variable	Description
1	Cost	3750.0	Cost	

Outbound Bilinks to Excel			
TreeAge Variable	Excel Cell	Excel Value	Description
Num_days	Num_days	15.0	
Per_diem	Per_diem	250.0	

BiLinks report for the tutorial example BiLink-after.trex.

23.2 Calculating payoffs using dynamic links: an example

This section describes how to create dynamic links to Excel. First open the Bilinks tutorial example, BiLink-before.tre and BiLink-worksheet.xls.

Once you have opened the tree and spreadsheet, there are three steps to setting up the dynamic link between them:

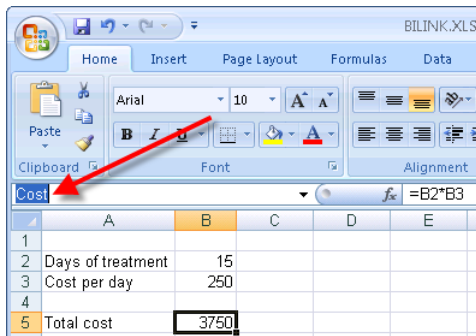
1. Link the tree to an output cell (or range of cells) in the spreadsheet.
2. Assign tree variable inputs to cells in the spreadsheet.
3. Define a payoff variable or other parameter in the tree to use the spreadsheet output.

23.2.1 Naming a spreadsheet cell

Before TreeAge Pro can connect to any input or output cells in the spreadsheet, these cells must be named. The Bilink tutorial example Excel file BiLink-worksheet.xls already has names assigned to the output cell (Cost) and the two input cells (Per_diem and Num_days).

To assign a name to a cell in Excel:

- Select the cell (or range of cells).
- Click in the Name box (see below) and type a one word name (like a TreeAge variable name).
- Press the *ENTER* key to save the name.



Name cell in Excel

If a cell (or range of cells) already has a name defined, it will appear in the Name box when the cell (or range of cells) is selected.

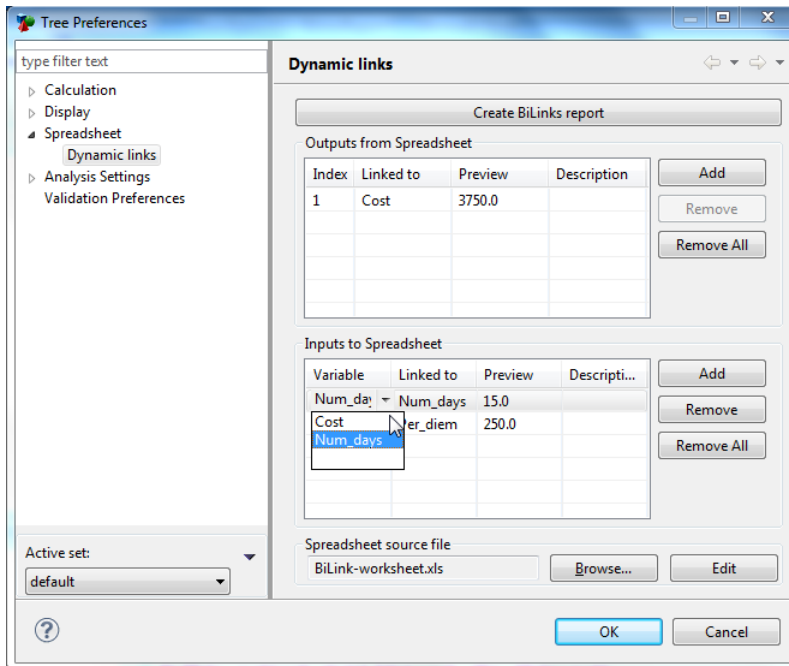
23.2.2 Linking to a spreadsheet cells

Once an output cell has been named, it can be linked to a TreeAge Pro tree as an input cell or an output cell (not both). In our model, we want to create two input cells for *Days of treatment* and *Cost per day* and one output cell for *Total cost*.

Both input and output cells are created via the Tree Preferences.

To create the dynamic links:

- Choose Tree > Tree Preferences from the menu or press the *F11* key to open the Tree Preferences dialog.
- Navigate to the category Spreadsheet > Dynamic Links.
- Click Browse to select the Excel spreadsheet (which must be saved).
- In the "Outputs from Spreadsheet" section, click the Add button. This should add the named cell *Cost*, which is the first named cell when sorted alphabetically.
- In the "Inputs to Spreadsheet" section, click the Add button. This should add the named cell *Cost*. Using the dropdown list within the Variable and Linked to columns, select *Num_days* in place of *Cost*.
- Repeat the above to create a link for the named cell *Per_diem*.



Tree Preferences - Dynamic Links. Note the dropdown menu for the Variable name.



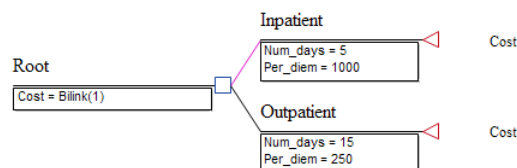
If you remove the path to the worksheet, TreeAge Pro will look for the worksheet in the same folder as the model. Use the Edit button to remove the path.

The links are now set up to send the Num_days and Per_diem values from TreeAge Pro to Excel, recalculate in Excel, and then receive the Cost value back. The inputs to Excel are already linked to TreeAge Pro variables in the Tree Preferences via the Linked to columns. We still need to link the output from Excel value Cost to our tree. This is done via the Bilink function using the index value listed in the "Outputs from Excel" section of the Dynamic Links Tree Preferences.

To define a TreeAge Pro variable using the Excel output cell:

- Right-click on the root node.
- Choose Define Variable > Cost from the context menu.
- Enter the definition *Bilink(1)* for the definition.

The argument 1 refers to the index value of 1 for the Cost dynamic link in the Tree Preferences.

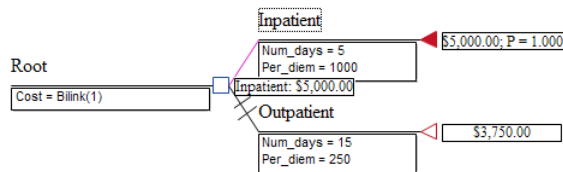


Tree using Bilink function

The inputs and outputs are now ready for calculation.

23.2.3 Calculate using dynamic links

Now, roll back the tree to verify that different values have been recalculated for each terminal node using the spreadsheet formula.



Bilink tree rolled back

Once the dynamic link is created, all calculations will utilize the linkage, not just roll back.

Excel must be running with the appropriate workbook open in order to create and/or use the dynamic links.

If the named linked cell is moved within the spreadsheet, the links will not be affected so as long as the linked cells' names are not changed or lost.



The Bilink tutorial example model BiLink-after.trex already has the appropriate links. However, they will not work on your computer until the path to the workbook is fixed in the Tree Preferences.

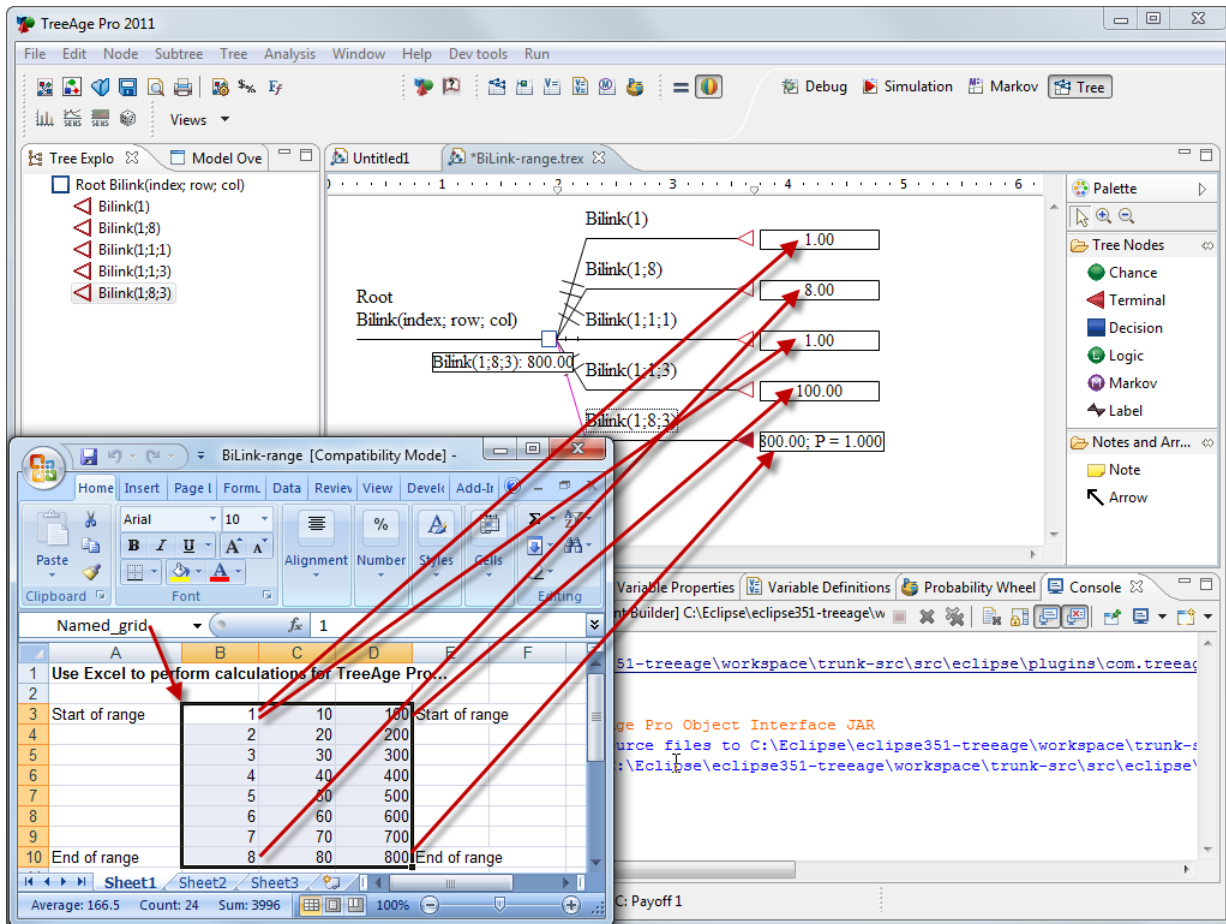
23.2.4 Using a single link to a range of cells

If a link is created to a named range of cells, the Bilink() function's index argument is followed by two more integer arguments — the row and column of the desired cell within the range. For example...

Bilink(1; 10; 2)

... will retrieve the value from row 10, column 2 of link #1's region of cells.

This feature is demonstrated in the Bilink tutorial example Bilink-range.trex and Bilink-range.xls. See below.



Bilink tree with range



The tutorial examples Excel model Bilink-range.trex will not work on your computer until the path to the workbook is fixed in the Tree Preferences.

23.2.5 Using Bilink() to create non-dynamic links

As mentioned above, if a dynamic link is set up without specifying any variables to input to Excel, then the link is not really dynamic. Use of TreeAge Pro's Bilink() function in this way is, in fact, the best way to create static links to cell values in an Excel spreadsheet.

The data is still pulled from Excel; however, no data is passed to Excel to possibly change the values of the data coming back.

23.2.6 BilinkName Function

Bilinks can also be referenced using the BilinkName function and the name of the bilink rather than its index. For example, in the Bilink tutorial example model Bilink-after.trex described earlier in this

chapter, the variable definition referencing the bilink could be entered as `Cost = BilinkName("Cost")` rather than `Cost = Bilink(1)`. The results would be the same.

24. Tools and Functions for Complex Trees

This chapter focuses on a number of features in TreeAge Pro that can be indispensable aids to both building and reviewing large and/or complex trees.

24.1 Working with very large trees

Many of the modeling exercises and examples in this manual are based on simple, small trees. The trees required in your own projects may be much more complex, perhaps including hundreds of variables, thousands of nodes, scores of distributions, etc. (Which is not to suggest that a bigger model is a better model, generally.)

Previous chapters detailed some of TreeAge Pro's many productivity features designed to make it easier to work with large trees:

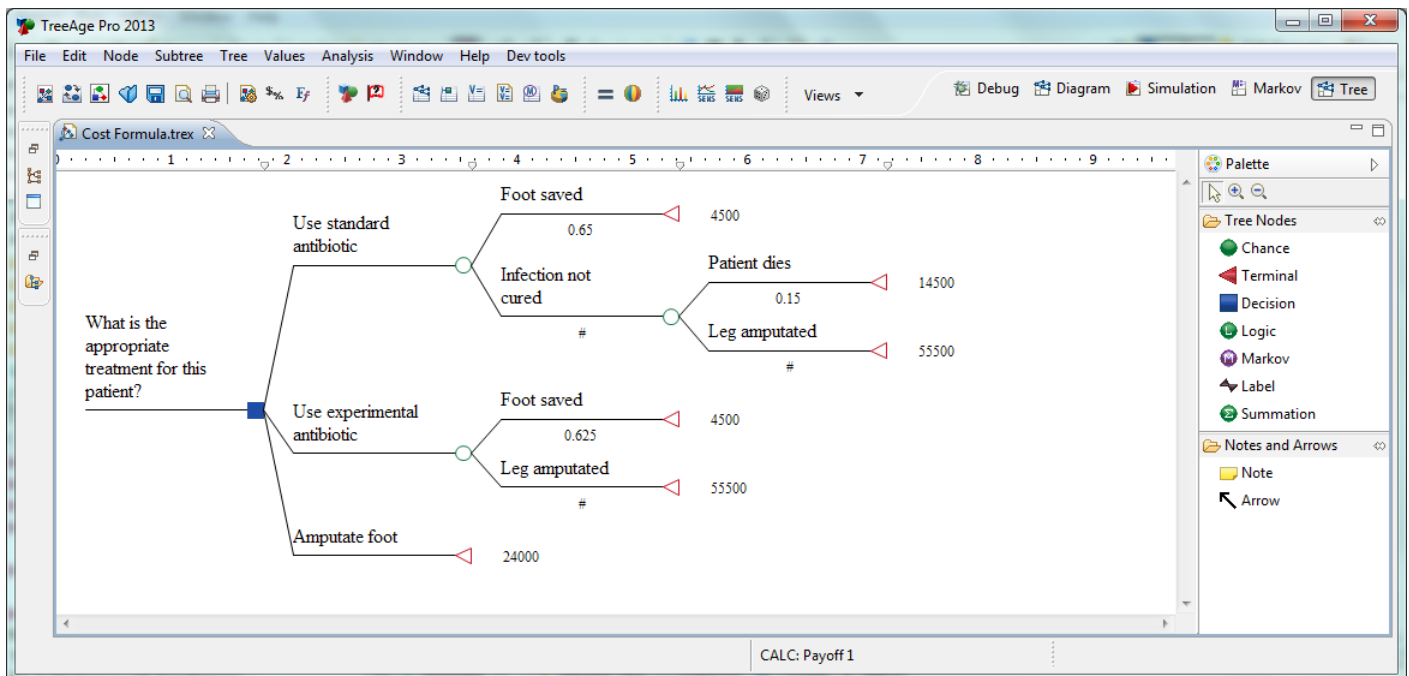
- Copy and paste subtrees.
- Collapse/hide subtrees.
- Use the zoom feature to see more/less of the tree structure in the editor.
- Use preferences that compress the tree.

This chapter focuses on additional productivity features relevant to complex modeling projects:

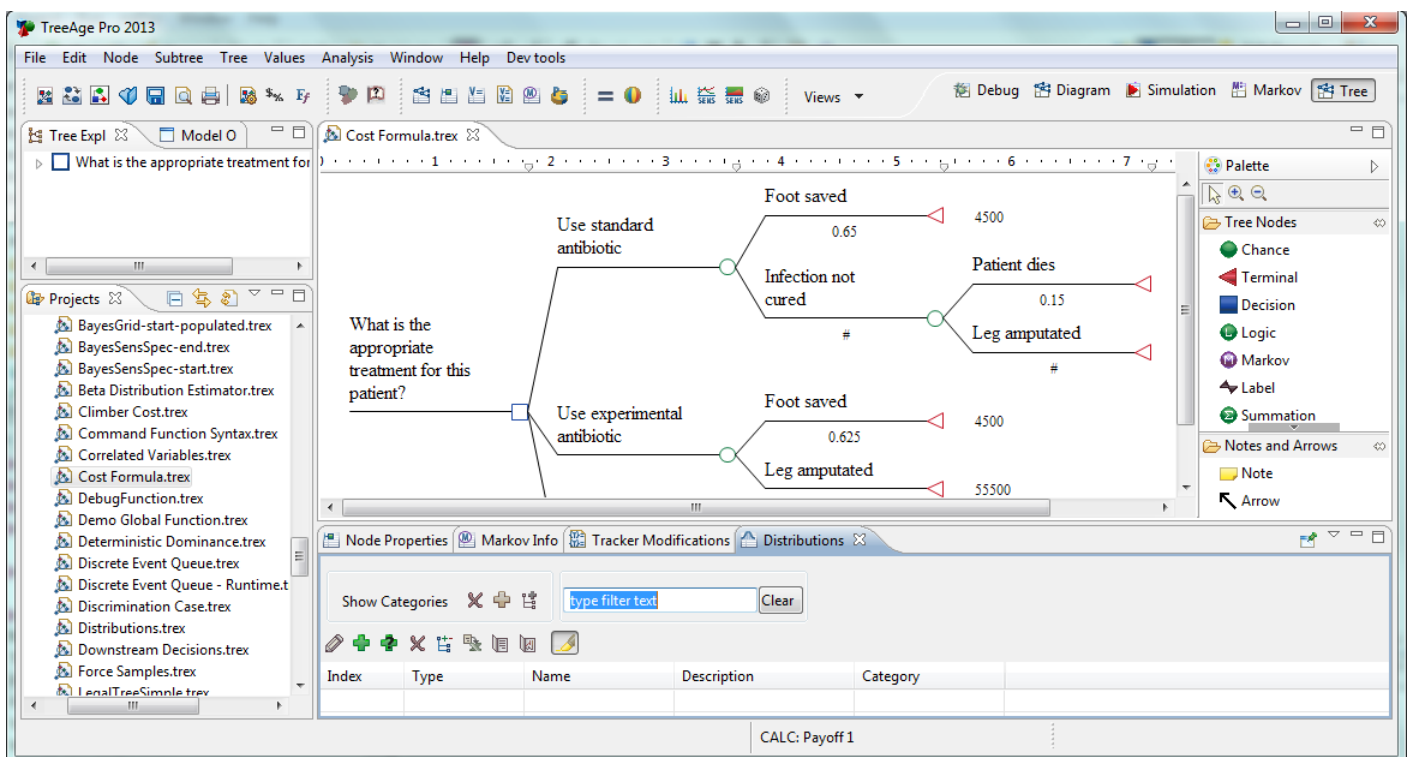
- Use the Explorer View and Model Overview for navigation
- View/edit the tree document's XML
- Clone subtrees (rather than copying/pasting)
- Creating user-defined functions with Python
- Link calculations to other trees/subtrees
- Using special functions to perform specialized tasks automatically during calculations (e.g., run a macro, enable debugging, or export the Global matrix)

24.2 TreeAge Pro Workspace, Tree Explorer and Model Overview

The TreeAge Pro Workspace is highly customizable, allowing you to move, resize, hide and maximize views and editors based on the task at hand. If you have a very large model, you might consider customizing the workspace in different ways. To rearrange your workspace for greater efficiency when working in complex models, you might try reducing one of the arrangements shown below.



Workspace with Tree Diagram Editor maximized



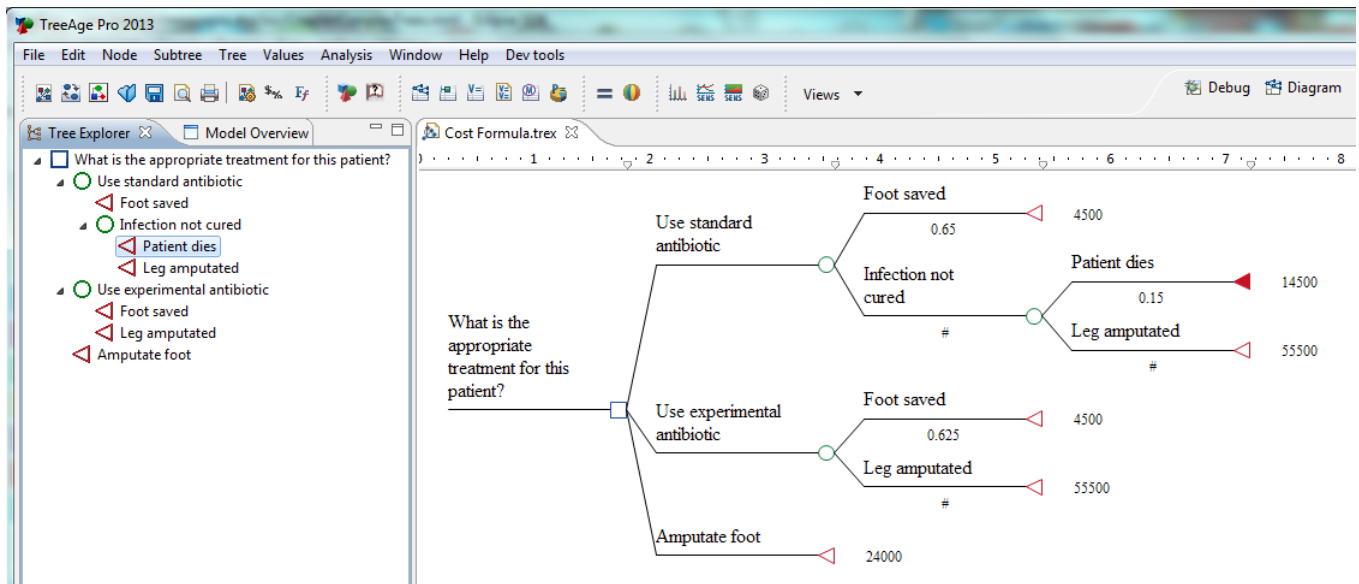
Workspace with Tree Properties View and Node Properties View visible

You can save these customized workspace presentations in Perspectives. Existing perspectives are shown at the top right corner of the workspace. Click the add button to create a new one.

Within the workspace, two views in particular, the Tree Explorer and Model Overview, can be especially helpful when working with large trees.

24.2.1 The Tree Explorer View

The *Tree Explorer* shows a collapsible text view of the model.

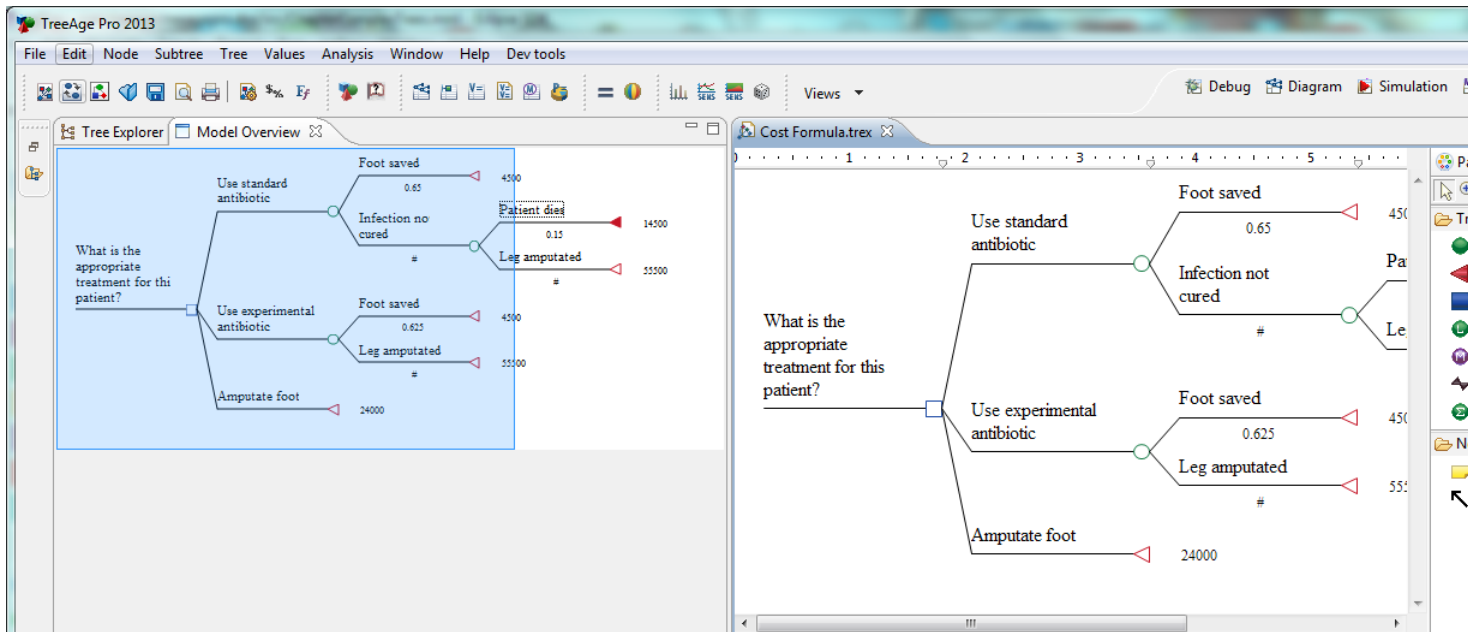


Tree Explorer

You can use the Tree Explorer to navigate to specific nodes. When you select a node in the Tree Explorer, focus is set on the same node in the Tree Diagram Editor. You can also open the node-specific context menu by right-clicking on a node in the Tree Explorer.

24.2.2 The Model Overview View

The *Model Overview* presents a high-level view of the overall model, highlighting the portion of the model that is currently visible in the Tree Diagram Editor.



Model Overview

Within the Model Overview, drag the highlighted area to a different area of the model to view a different area of the model in the Tree Diagram Editor.

24.3 Viewing/editing document XML

TreeAge Pro documents are stored in XML format. TreeAge Pro provides the option to edit the document's XML in an XML editor or a text editor.



Unless you are quite familiar with the XML syntax used by TreeAge models, it is recommended that you use the TreeDiagram Editor to make all edits in models (in particular, adding/deleting nodes), rather than editing the XML text file itself. It is easy to render a model unreadable by making bad edits to the file XML.

However, it is simple to manually create copies/backups of *.TREX files before editing the XML. The Projects explorer view also supports robust *text compare/local history* features, via the right-click menu. This makes it possible to track and remove/reverse selected edits to any of your project files (whether changes are made in the Tree Diagram editor or a text/XML editor). Finally, the powerful Search/Find/Replace functionality used in combination with the Project explorer view allows *regular expression* search/replace within the XML (and any other text-based project documents).

To open a tree in an XML editor:

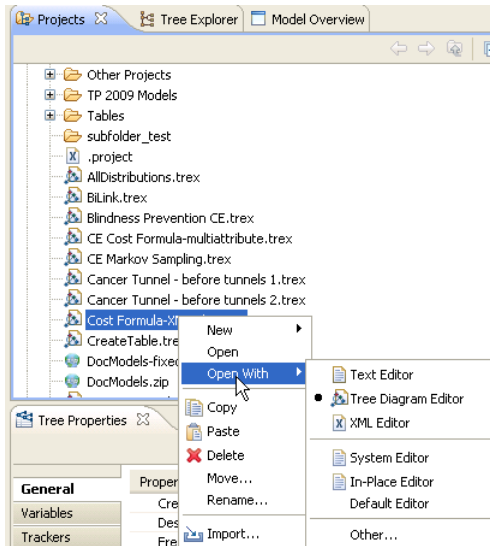
- Right-click on a model in the Projects explorer.
- Choose Open With > Text Editor.



The "default" method of opening the selected model will change when opened in this way. This means that next time you simply double-click on the file, it will again use the "Text Editor" to open. Therefore, it is necessary to right-click and choose Open With > Tree Diagram Editor in order to restore the original, default opening method.

To open a tree in an XML editor:

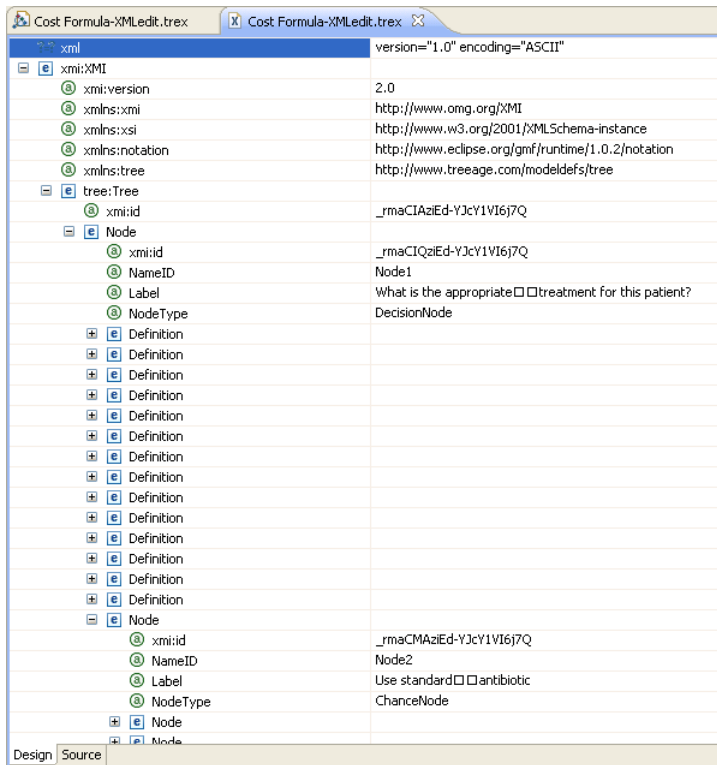
- Right-click on a model in the Projects Explorer.
- Choose Open With > XML Editor.



Open with XML Editor

The tree will appear in an XML editor within the workspace. You can also open the tree in a Text Editor, which shows the XML in plain text format.

The Special Features tutorial example tree "Cost Formula" is shown below in the XML editor.



XML Editor

XML consists of a hierarchical structure of textual data. In the XML editor, each level of the structure can be expanded or collapsed.

A portion of the basic XML structure of a TreeAge Pro tree is presented below.

```
Document
- tree:Tree
- - Node
- - - Definition
- - - Node (child)
- - - - Node (child)
- - Variable
- - - Sensitivity Range
- - Categories
- - Preferences
- notation:Diagram
```

XML structure

Some notes on the XML format:

- The highest-level elements within the tree document structure are tree and notation.
- The *tree* element includes the building blocks of a model - variables, nodes, preferences. etc.

- The *notation* element contains details related to presenting the model in the Tree Diagram Editor.
- *Variable* elements are nested within the *tree* element because variables exist at the tree level.
- *Definition* elements are nested within a specific *Node* element because variable definitions are defined at the node-level.
- Some XML elements are not presented above (Markov info, payoffs, etc.), but they do exist within the XML structure.

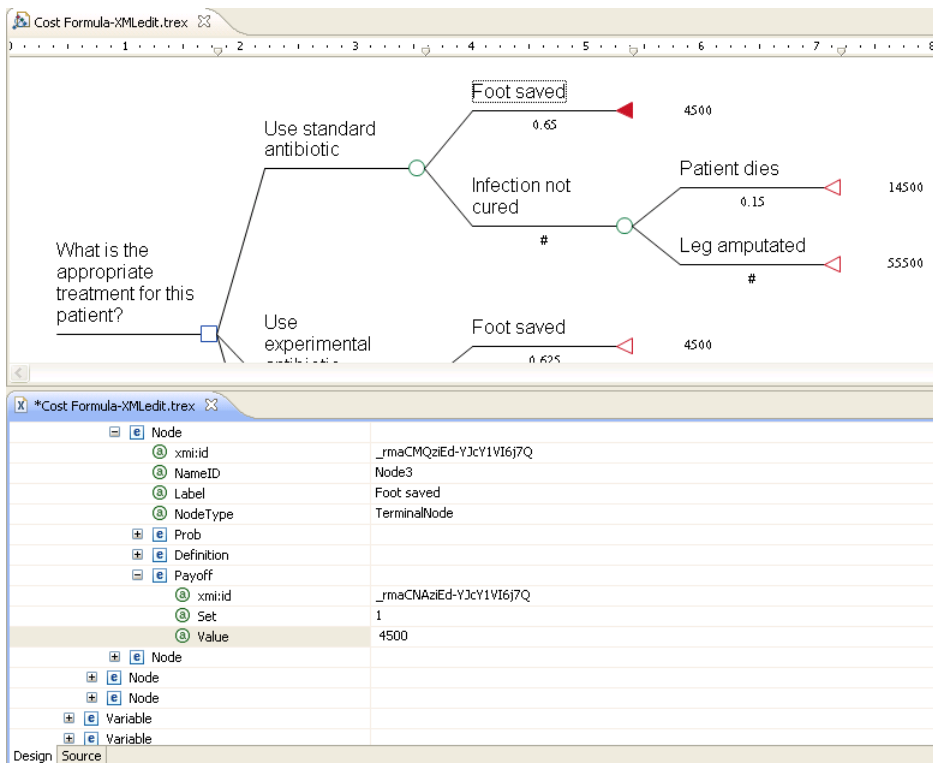
A more complete, but not exhaustive, list of XML elements is presented in the following table:

Element	Description	Parent Element
tree	Highest level of the tree structure.	<top level>
notation	Controls presentation of the model in an editor.	<top level>
Variable	Variable name and properties.	tree
Sensitivity Range	Sensitivity analysis range for a variable.	Variable
Node	Node and its properties. Nested according to its location within the tree.	tree or Node
Definition	Variable definition associated with a node.	Node
CategoriesRoot	Variable categories for the tree.	tree
PreferenceSet	Tree preferences.	tree
Termination	Termination condition for a payoff set/calc method	Node (type Markov)
Prob	Branch probability.	Node (with parent chance node)
MarkovData (state)	Collection for Markov information associated with Markov State (state rewards).	Node (with parent Markov node)
MarkovData (transition)	Collection for Markov information associated with Markov Transition (transition rewards, jump state).	Node (in transition subtree)
Payoff	Payoff expression for a specific payoff set at that terminal node	Node (type Terminal)

XML Elements

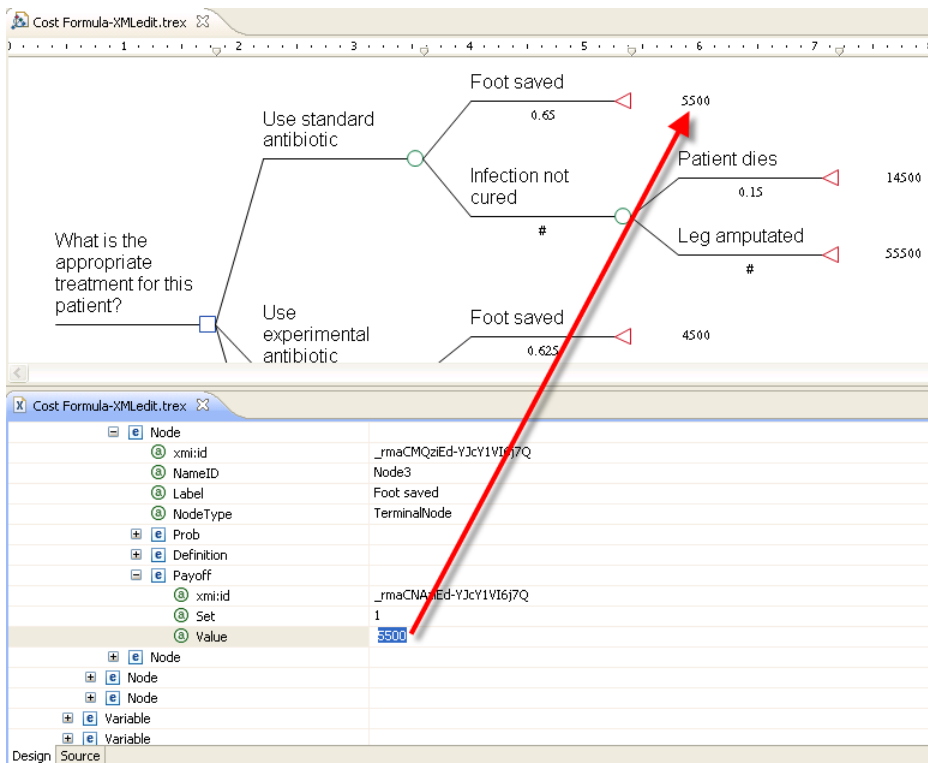
24.3.1 Editing XML data

You can edit the structure of the model or properties of a model element within the XML view of the document (although this is not recommended). Consider the following image of a tree both in the Tree Diagram Editor and the XML Editor.



Tree Diagram Editor and XML Editor

Consider the *Value* attribute within the *Payoff* element for the *Node Foot Saved*. If I change the Value from 4500 to 5500, then save the file in the XML editor, the Tree Diagram Editor will reflect the same change.



Edit model via XML Editor



Be sure to save all changes in the Tree Diagram Editor before opening the XML Editor for changes. Otherwise, you run the risk of overlaying your editor changes with a freshly updated version from the XML Editor.

Additional notes on XML editing:

- You can delete XML elements or move elements around within the XML structure.
- You can add new elements to the XML structure, but the `xml:id` must have a unique value for the new item.
- The File Search (*Control + F*) results are listed in XML format, so some knowledge of the XML format can be useful.

24.4 Cloning subtrees

In addition to being able to duplicate existing subtrees by copying and pasting, it is also possible to create *clones* of subtrees. The basic difference between attaching clones and pasting copies is that a copied/pasted subtree can be edited; these copies do not automatically update when changes are made to the original. Clone copies, on the other hand, are linked dynamically to the original *master* subtree, and are not directly editable. A clone copy always automatically takes its structure and other contents from its clone master (unless the relationship is explicitly broken).

Clone copies function exactly like their associated clone master. However, variables within the clone master (and copy) can reference different variable definitions. This allows you to use different values for those variables within the clone master and copy. See cloning example below.

A few notes on clones:

- Multiple clone masters can be used in a tree, and can be nested.
- Variables (and state bindings) *used* in a clone master can *calculate differently* in clone copies if the variables are *defined* outside the master/clone.
- Cloning only works within a single tree.
- A clone master copied as part of a larger subtree will *not* be pasted as a clone master (or as a clone copy). A clone copy, when copied as part of a larger subtree, will be pasted as a regular subtree, not a clone copy.

24.4.1 Creating clone masters and attaching copies

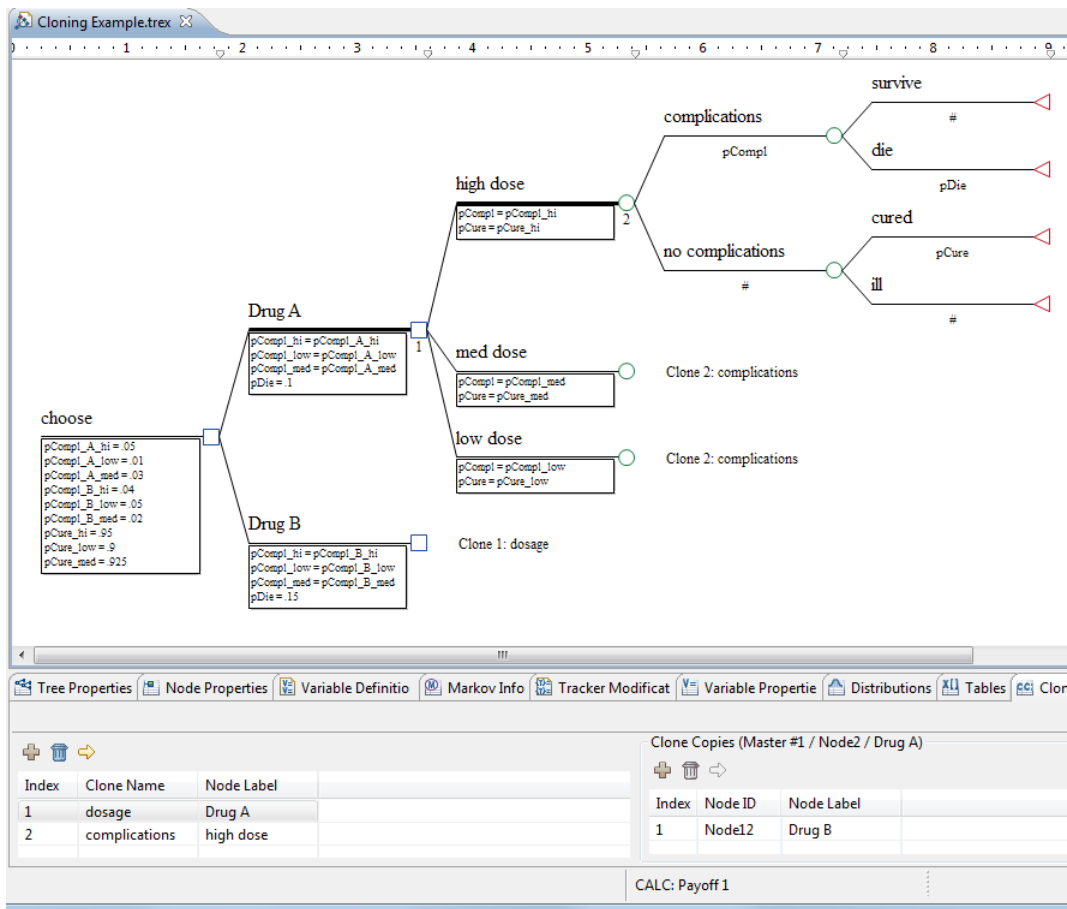
The first step in the cloning process is to select a clone *master* subtree for the purpose of being replicated at another location in the tree. The clone master does not have to be a finished structure; all that is required is a single branch. The master subtree can continue to be updated after copies of it have been attached to other nodes in the tree.



In addition to the standard method described below for creating/attaching/detaching clones, TreeAge Pro also supports handling most of these operations in the Clone Master/Copies view as shown here. This view lists the tree's clone masters and all of the clone copies for a selected clone master. See the end of the section for more details.

Each subtree that is a designated clone master can be identified by a heavy bar beneath the branch leading to its root node. Like a copied subtree, what will be "cloned" will not include the root master node - only its subtree.

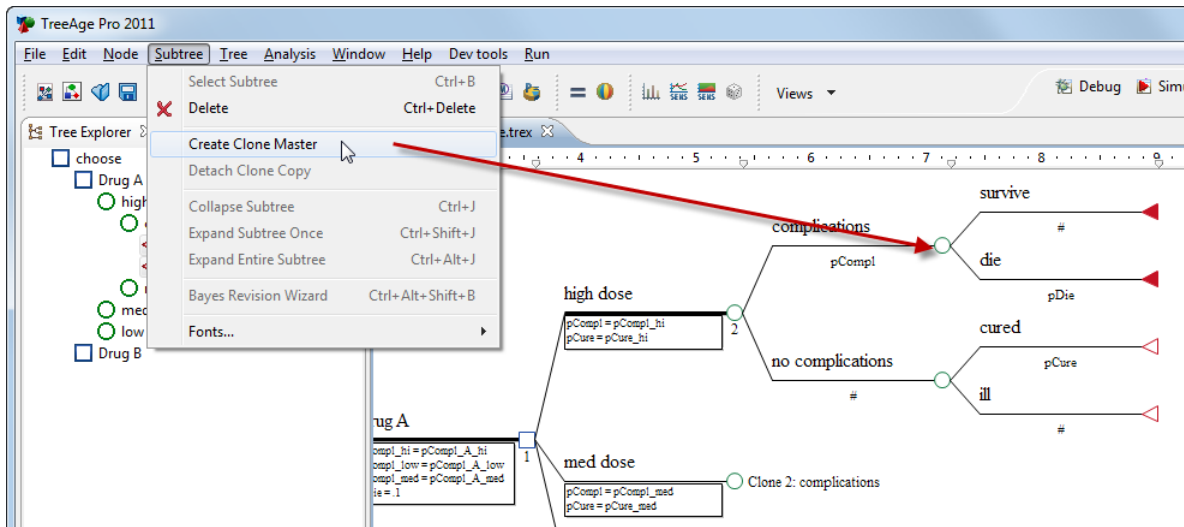
The remainder of this section uses the Health Care tutorial example model, Cloning Example.trex, shown below with existing clone masters/copies listed in the Clone Master/Copies View.



Cloning Example tree

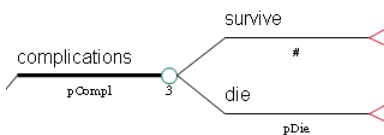
To create a clone master:

- Identify the master subtree and select it.
- Choose Subtree > Create Clone Master from the menu.
- Provide a short, descriptive name to identify this clone master (survival).



Create clone master

Clone masters are also assigned a numeric index which appears next to the clone master node.



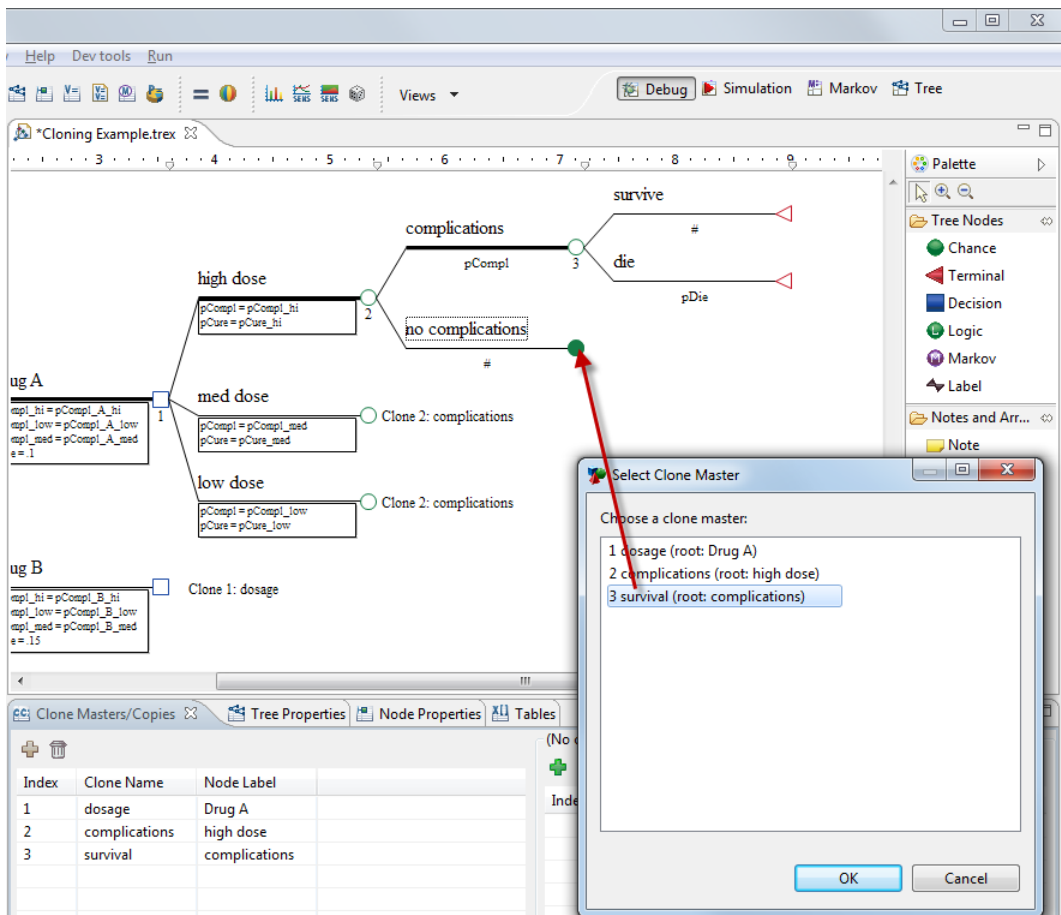
Clone master identified by heavy bar and index number

Once the clone master has been created, the next step is to attach the clone copies at the appropriate node(s).

Clone copies can only be attached to nodes which have no branches. The nodes where you attach a clone copy should have the *same node type* as the root node of the master subtree. Attaching a clone copy to a node will not automatically change the type of the node to match the clone master's root node. This change can be made manually, if necessary, either before or after attaching the clone copy. In our example, we will attach the new clone master to the *no complications* node (note that its subtree has been deleted before the attach clone steps below).

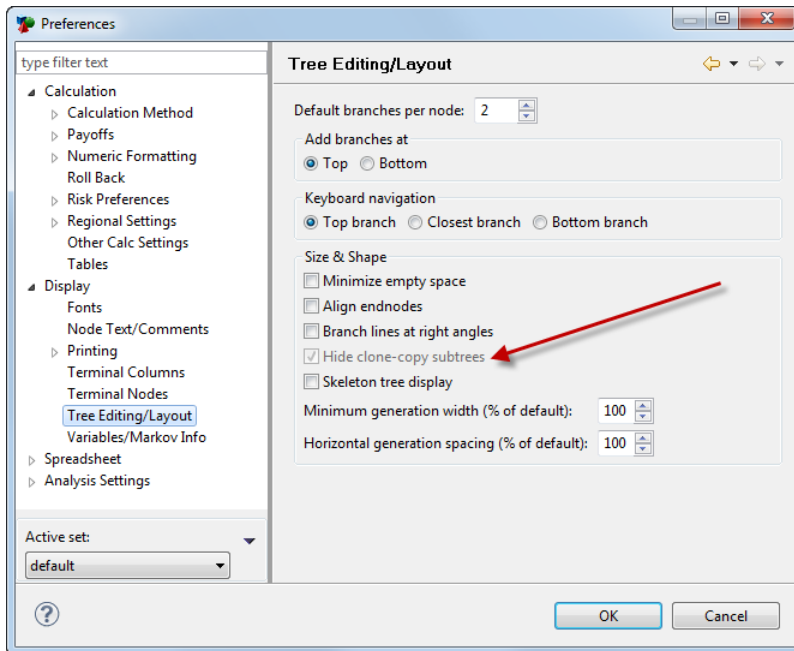
To attach a clone copy:

- Select an appropriate node (i.e., a node with no descendants).
- Choose Subtree > Attach Clone.
- If you have only a single clone master in the tree, it will be attached to the selected node automatically. Otherwise, you must select the name of the appropriate clone master from a list, and click OK.



Attach clone copy

Currently, clone copies are always hidden, showing only references to the clone masters. In the future, the "Hide clone-copy subtrees" option in the Tree Preferences will be enabled so that you can uncheck the option. See below.



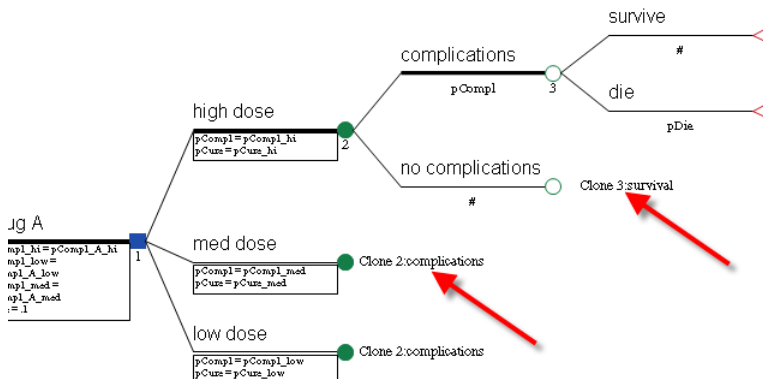
Tree Preferences - Show/hide clone copies

The "Hide clone-copy subtrees" option is currently checked and disabled. In the future, the option will be enabled to allow you to uncheck it to see the entire clone copy subtrees, which will be done in gray in order to distinguish them from editable subtrees.

To see a clone copy displayed prior to this tree preference being fixed:

- Save a temporary copy of the model.
- Detach the clone copy.
- When prompted, choose to create an editable copy of the clone master.

Currently, with the "Hide clone-copy subtrees" option always checked, you see a reference to the clone copy in the tree, but not the entire subtree. This reduces the overall size of the displayed tree. Showing/hiding clone copies does not affect calculations in any way.



Clone copies hidden

Reducing the tree size by hiding clone copies enhances the clarity of the tree. The essential features of the replicated subtree can be seen by examining the clone master, which is the only instance of the subtree being displayed. In addition, the cloning linkages within the model, which might otherwise be missed, are clearly visible as each clone copy indicates the master to which it is linked to.

24.4.2 Destroying and detaching clones

If you want to remove clone copies from a tree, you can either destroy the current clone master (automatically detach its clone copies) or detach the selected clone copies one at a time.

There are two ways to destroy a clone master.

To eliminate a clone master in the tree:

- Select the clone master subtree or the subtree's root node.
- Choose Subtree > Destroy Clone Master from the menu.

To eliminate a clone master using the Clones dialog:

- Select the clone master in the list, and click the Delete ("Trashcan") toolbar button.

Destroying a clone master will “un-publish” the subtree. The subtree which was formerly the clone master will remain in the tree; all of its clone copies will be removed. You will be prompted with the option to replace those clone copies with independent copies of the clone master.

To detach clone copies:

- Select the clone copy subtree or the subtree's root node.
- Choose Subtree > Detach Clone Copy from the menu.

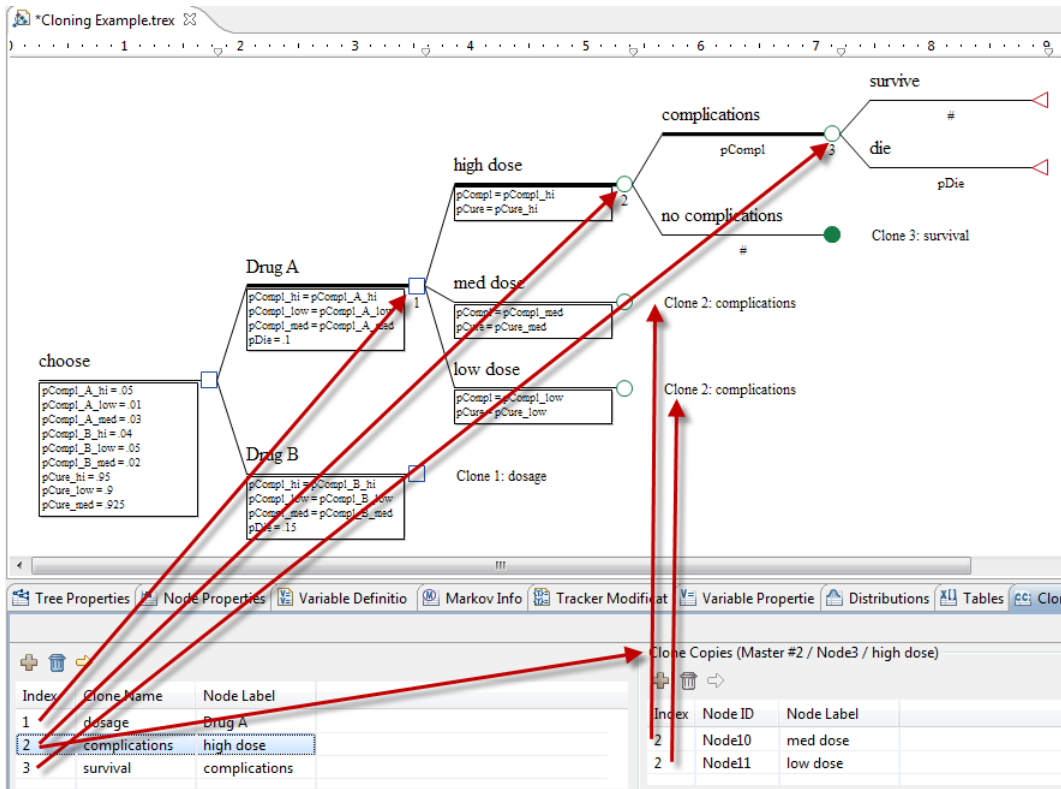
You will be asked if you want to keep an editable copy of the cloned subtree at the selected node. If you answer Yes, then the clone copy subtree is replaced by an independent copy of the clone master subtree. If you answer no, the clone copy subtree is simply removed.

24.4.3 The Clone Masters/Copies View

The Clone Masters/Copies View allows you to view and edit clones within the tree.

To open the Clone Masters/Copies View:

- Choose Views > Clone Masters/Copies from the toolbar.



Clone Masters/Copies View

The arrows in the figure above demonstrate the relationships between the clone masters/copies in the view and in the model.

Clone masters are listed in the left pane of the view. When one is selected, the clone copies associated with that view are listed in the right pane.

Both clone masters and clone copies can be deleted through this view with the same options and restrictions associated with deleting these elements in the Tree Diagram Editor, as described in an earlier section.

The arrows in the two toolbar of the Clones View can be used to move directly to either a clone master or a clone copy in the Tree Diagram Editor.

24.4.4 Nesting Clones

An important aspect of clones is the ability to nest clone masters. In other words, a single subtree may be comprised of multiple independent clone masters and also include various clone copies.

The tree shown below in the next section has three nested clone master subtrees. Clone masters 2 and 3 are nested within other clone masters and their copies.

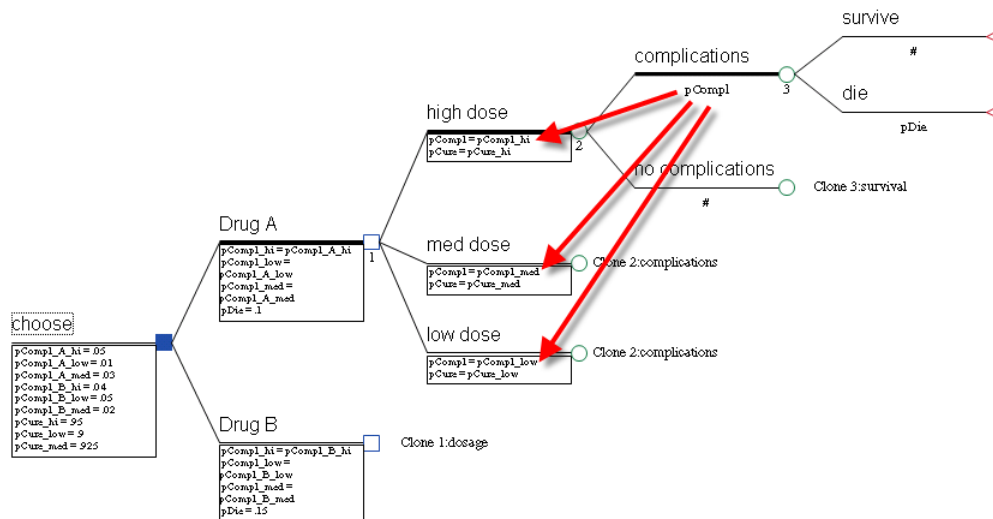
You cannot attach a copy of a master subtree to itself. To create recursion, use a Markov node instead.

24.4.5 Using variables within clones

Calculations in clone copies are performed as if a full copy of the master subtree existed at the location of the clone copy. This is true even if you have turned off the display of clone copies in the tree.

Since clone copies are identical to the clone master, *variables* must be used in the clone master if you want its copies to have *different probabilities and values*. The probability, reward, and payoff expressions in the clone master and its copies will all use the *same variables*. However, each will be able to use *different values for these variables* based on how the variables are defined outside the clones. Variables referenced within the clones can be defined outside the clones. By defining variables differently to the left of the clone master and its clone copies, different values can be used by the same variable references within the clone master and copies.

The Health Care tutorial example tree "Cloning Example" provides a relatively simple example of using variables with clones.



Clones referencing a variable with different definitions

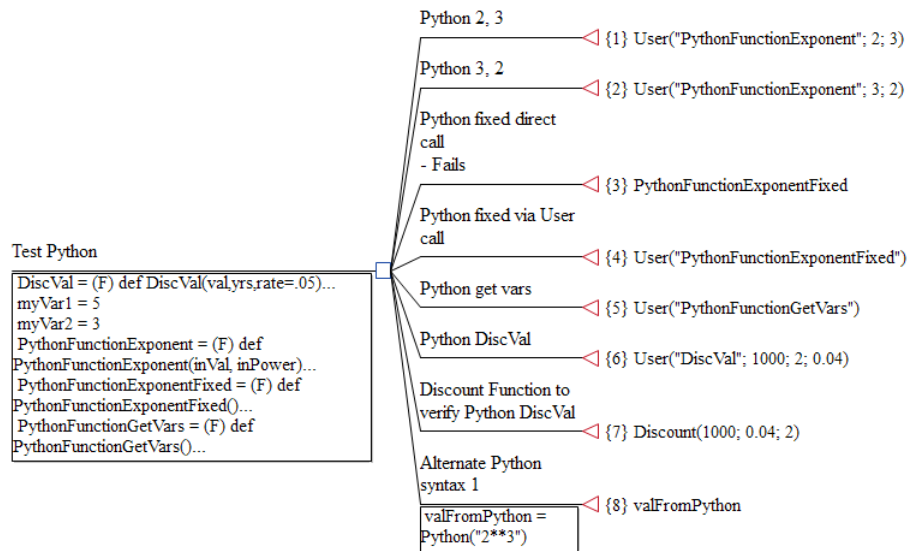
Note that the variable *pCompl* is used in the clone master, meaning it is also used in the clone copies. However, at the root of the clone master and clone copies, the variable is defined differently. This allows a different probability to be used for each of the subtrees.

24.5 User-defined Python functions

If there is no built-in function that does what you require, you can write a user-defined function using Python script and simply embed it in a regular variable definition in your tree.

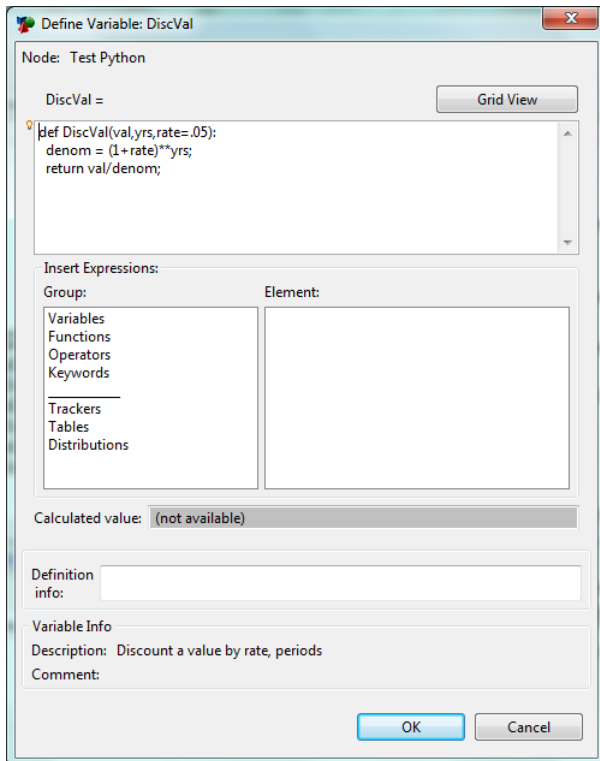
24.5.1 User-defined Python function variable and syntax

The Special Features tutorial example model, Python Example, contains the functions discussed in this section.



Python Example tree

User-defined Python functions are created within a variable definition. For example, a variable called DiscVal could be added to the tree. Then DiscVal could be defined at a node (probably the root node) with the following function entered as its definition:



Python function in variable definition

The function declaration is the first line:

```
def DiscVal(val,yrs,rate=.05):
```

After the Python keyword *def*, repeat the name of the tree variable using the same case (Python is case sensitive). After the function name, parentheses are required. Arguments are separated by commas; an optional argument list can be entered within the parentheses and optional arguments, starting at the right, can have default values. The declaration ends with a “.” (colon) and a carriage return.

Lines following the declaration must use tabs or spaces to express indent levels; this is how Python interprets blocks of code. Statement lines are optionally ended/separated with a semicolon and use “#” at the start of a comment line. The *return* statement is used to return a value from the function; the end of the function is implicit.



Tabs cannot be entered into the variable definition Formula Editor; enter a consistent number of spaces instead to represent the nesting level for a block of code within the Python function. Python functions can be edited in a separate editor which supports tabs. Then the contents can be pasted into the variable definition Formula editor.



For more information on the Python language and syntax, go to <http://www.python.org/>.

In the tree view, user-defined function definitions are prefixed with “(F)”. Only the declaration line will be shown unless wrapping is turned on under Variables Display preferences.

24.5.2 Calling a user defined Python function

User-defined functions can be used in a payoff, probability, or any other expression in the tree. If your user-defined function requires arguments, then it must be called via the `User()` function:

```
User("Python function"; argument 1; argument 2; ...)
```

You might use the `DiscVal` function defined in the previous section in a variable definition as follows:

```
cRx = User("DiscVal"; 5000; _stage; rate)
```

The example tree's payoffs 1, 2, 4, 5 and 6 use this format, although a couple of them require no arguments.

If no arguments are required for a particular user-defined function, then simply call the function using its variable name without parentheses or a `User()` call.

For example, the following function requires no arguments:

```
def PythonNoArgs():
```

Therefore, it can be referenced directly by name as follows:

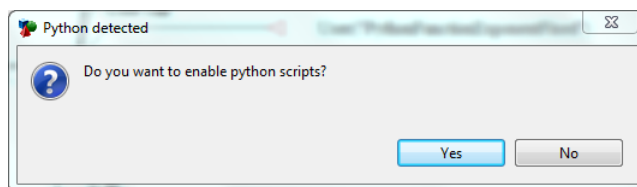
```
myVar = PythonNoArgs
```

The example tree's payoff 3 uses this format.

The example tree's payoff 7 simply checks to ensure that the Python discount function returns the same value as the built-in `Discount` function. The example tree's payoff 8 references a variable that is defined using the `Python("Expression")` format described later in this chapter.

24.5.3 Script security

When you open a tree with a user-defined Python function, you will be prompted to enable Python scripts.



Enable Python scripts dialog

The types of Python commands useful in trees are not a security concern, but the core language does have file and Internet access methods which could theoretically be misused. The prompt is intended to encourage examination of scripts in models from unknown sources (e.g., search for “def” and “python”).

24.5.4 Using the "treeage" object

Although values (e.g., of variables) can be passed to the Python user-defined function via a list of arguments, the script can also evaluate tree variables, trackers, and other expressions directly:

```
def mySpecialFunction():
    ct = treeage.eval("trackAEs");
```

In the context of the Python function, *ct* would be set equal to the value of the expression *trackAEs* in the current node context of the tree – e.g., at the node whose calculations reference *mySpecialFunction*.

The `.eval()` method is only one of numerous methods available via the “treeage” object.

Treeage object function	Description
<code>treeage.eval("expression")</code>	Returns the value of <i>expression</i> from the tree.
<code>treeage.debug("title", "msg")</code>	Output to trace console view for tree (if detailed debugging preference is on).
<code>pt = treeage.getParallelTrials()</code>	Reference to parallel trials info.
<code>matrix = treeage.getGlobalMatrixN(n)</code>	Sets a reference to one of the global matrices, whose data can then be accessed/changed. See following set of rows.
<code>matrix.getRows()</code> <code>matrix.getCols()</code>	Get the current number of rows, columns in the matrix.
<code>matrix.getElement(int row, int col)</code>	Get the value from a cell in the matrix at row <i>row</i> , column <i>col</i> .
<code>matrix.setElement(int row, int col, double val)</code>	Set the value of a cell in the matrix to <i>val</i> at row <i>row</i> , column <i>col</i> .
<code>matrix.incrElement(int r, int c, double val)</code>	Increment the value of cell in the matrix by the amount <i>val</i> .
<code>matrix.fill(double val)</code> <code>matrix.fillCol(double val, int col)</code> <code>matrix.fillRow(double val, int col)</code> <code>matrix.fillCol(double val, int col, int rowsUpperLimit)</code> <code>matrix.fillRow(double val, int row, int columnsUpperLimit)</code>	Populate the entire matrix with the <i>val</i> provided. Populate all cells in column <i>col</i> with the <i>val</i> provided. Populate all cells in row <i>row</i> with the <i>val</i> provided. Populate the cells in column <i>col</i> with the <i>val</i> provided from row 1 to row <i>rowsUpperLimit</i> . Populate the cells in row <i>row</i> with the <i>val</i> provided from column 1 to column <i>colsUpperLimit</i> .
<code>matrix.getIndex()</code>	Returns the index of the matrix.
<code>matrix.clear()</code>	Empties the matrix of all data.

Python treeage object syntax

24.5.5 Performance of the TreeAge Python interpreter

If performance is critical in your model, try to use the Python interpreter efficiently. For example, a `treeage.eval()` call is slower than passing a variable value as a `User()` function argument. Because

of the Python's global interpreter lock, inefficient scripts may be exacerbated during multi-threaded simulations.

24.5.6 Why TreeAge Pro uses embedded Python

Python is an easy-to-use object-oriented programming and scripting language which includes standard elements like looping, type conversion, etc.:

```
for i in range(1 to 20):
    s = str(i);
    ct=ct+treeage.eval("ctEvt"+s);
```

Your user-defined functions can also import from Python's powerful set of built-in modules as well as many open source, third-party modules.

```
from random import log10;
return math.log10( val );
```

To support user-defined functions, the TreeAge Pro installation includes the core Python modules only. To test Python scripts outside of a TreeAge model (or if you require custom modules), simply download and install Python. This will enable you to use IDLE or a Python command-line to check syntax.

To download the full Python language, go to <http://www.python.org/>.

A good reference guide to the Python language is *Python in a Nutshell, 2nd Ed.* (O'Reilly, 2006).



The "treeage" object is not available *outside* TreeAge Pro when using an external Python IDE.

24.5.7 Other Python syntax options

TreeAge Pro supports several syntax options for calling Python functions. The standard ones previously mentioned as well as others are described in the following table.

Function Syntax	Description
<i>myPythonFunction</i>	Call Python functions that require no arguments by simply referencing the function name.
<i>User("myPythonFunction"; argument1 ; ...)</i>	User command to call Python function and pass arguments to the function.
<i>Python("expression")</i>	If the current Python can successfully evaluate the variable name, expression, or function, the Python() function will return that value. If it cannot evaluate it, an error is returned. The Python() function returns a numeric value. Example: <i>Python("10+1")</i> returns 11 Example: <i>Python("int('0xff', 16)")</i> returns 255

Function Syntax	Description
<i>The Python syntax below is not yet supported in TreeAge Pro</i>	Not yet available.
<i>PythonFunc("module"; "function"; argument1 ; ...)</i>	Execute a function from an imported module. This will only work if the module has already been imported. Example: <i>PythonFunc("random"; "normalvariate"; mu; sd)</i> returns a random number from Normal(mu,sd) where <i>mu</i> and <i>sd</i> are numbers
<i>PythonGetList("list"; "treeage table")</i>	The PythonGetList() function copies a two-dimensional Python nested list into a TreeAge table. The Python expression can be a literal list (as seen below), a variable containing a list, or a function returning a list. Example: <i>PythonGetList("[[0,1,.2],[1,.15,.25]]" ; "myTable")</i> loads myTable and returns the number of rows (2)
<i>PythonRun("command"; ...)</i>	Returns 0 for failure or 1 for success. Specify a valid Python statement. To run a series of scripts or modules, separate multiple double-quoted paths using semicolons. If Python successfully runs the last specified statement, the PythonRun() function will return a value of 1. If Python encounters an error evaluating your commands, the PythonRun() function returns a 0 value -- not an error. To force a TreeAge Pro error, use the result as the denominator of a division operation. To return the value of variables or functions created or modified by the executed commands, use the Python() function. Example: <i>PythonRun("from random import *"; "seed(1)"; "x=random()") = 1</i> <i>value = Python("x") = .134364...</i>
<i>PythonRunFile("scriptfile" ; ...)</i>	Returns 0 for failure or 1 for success. Specify the full path and filename of a valid Python script text file or module to run; or, specify just the file name if it is stored in the Python default path. To run a series of scripts or modules, separate multiple double-quoted paths using semicolons. (Note that forward slashes can be used instead of backslashes in the path specification.) Example: <i>PythonFile("C:\Python24\lib\TransplantQueue.py")</i> Using PythonRunFile() to execute a file is equivalent to having Python evaluate the execfile() function: <i>PythonRun("execfile('C:\Python24\lib\TransplantQueue.py')")</i> If the script/module executes successfully, use the Python() function to return the value of variables:

Function Syntax	Description
	<code>queueWait = Python("now()")</code>

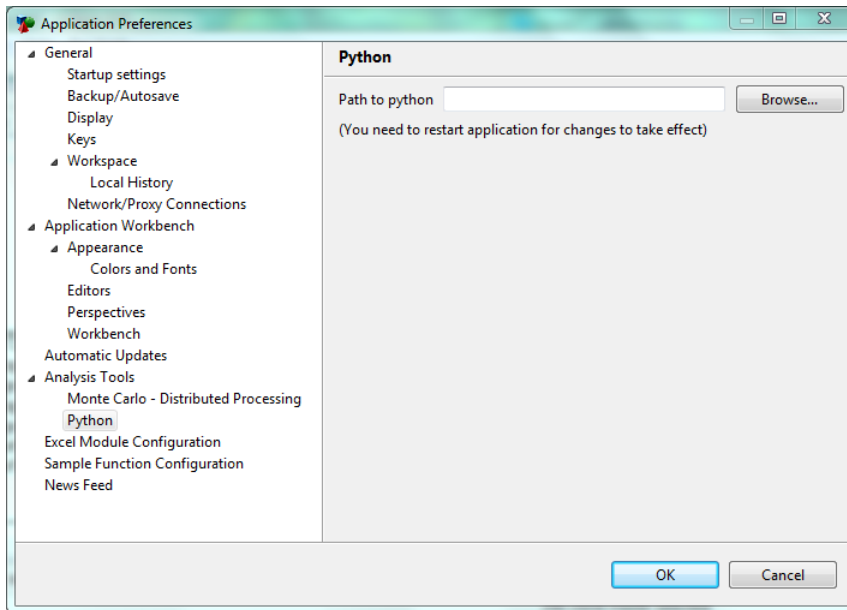
Python syntax options

24.5.8 Overriding the Python interpreter

Python version 2.5.1 is embedded in TreeAge Pro. However, you can choose to install Python directly onto your computer - perhaps to use a later version of Python or to import additional Python modules outside the base version. You can then direct TreeAge Pro to use the Python installation instead of the embedded version.

To redirect the Python interpreter:

- Choose Window > Application Preferences to open the Preferences dialog.
- Select the category Analysis Tools > Python.
- Enter or browse to the path where the Python executable is installed on your computer.



Preferences - Python

24.6 Node(), Tree(), User(), Global() and other special functions

TreeAge Pro includes a variety of special functions which can be useful in complex models; several of these functions use string arguments. These functions are described in the next several sections of this chapter.

24.6.1 String-arguments and concatenation

Most TreeAge Pro functions accept only numeric arguments, although those numeric arguments can be provided via variables and expressions. However, a variety of functions use text/string function arguments. For example, the Tree function requires string arguments.

```
Tree("myTree_1.tre";"Node(1;0)")
```

Most string arguments can be constructed using concatenation; in other words, text and calculated values can be combined into a new string. For example, the filename in the Tree() function could be concatenated (with a variable called "pick", in this case):

```
Tree("myTree_"+pick+".tre";"Node(1;0)")
```

Similarly, the worksheet name in the function...

```
Command("Excel";"ExportGlobalMatrix";Workbook;SheetName;CellName;LabelText)
```

... could be concatenated with a variable:

```
Command("Excel"; "ExportGlobalMatrix";"Outfile.xls";"sheet_"+_trial+"_ "; ...)
```

24.7 The Node() function

The Node() function can act as an internal linking function within a particular tree; this allows nodes in different parts of a tree to communicate. Some possible uses for the Node() function include:

- Access calculated expected values in non-active payoff sets (in terminal columns only).
- Reuse a subtree at different locations in a tree for the active payoff set (similar to the way cloning is used, but with greater flexibility).



Note that the Node() function can be called from within another Node() function.

The simplest form of the function is presented below...

```
Node( attribute )
```

...where *attribute* is an integer referring to a payoff set. This value must be an integer referring to an enabled (not necessarily active) payoff set. Calculations are performed at the node where the function is called.

The Terminal Columns model described in the Tree Display Preferences and Options Chapter uses this Node() function syntax to display the expected value from a non-active payoff set in a terminal column.

The more complete function syntax is presented below:


```
Node( attribute; method; branch list )
```

The argument *attribute* refers to the payoff set to use for the calculation.

The argument *method* refers to the type of calculation to perform.

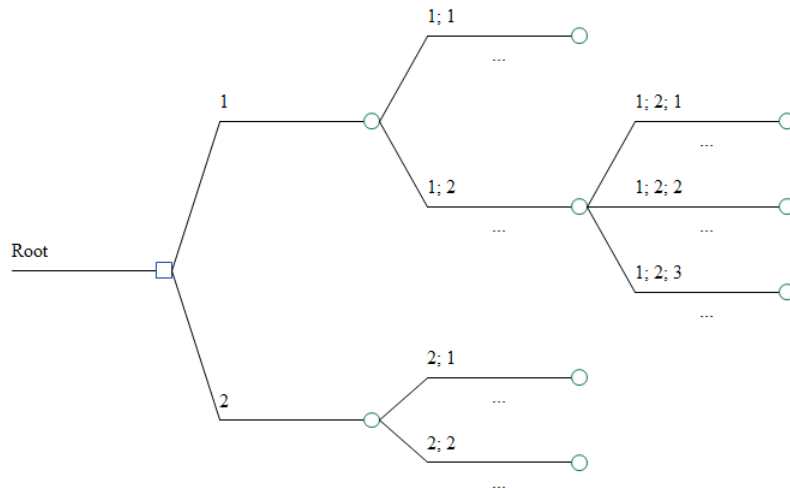
The argument *branch list* refers to the node at which to perform the calculation.

The different forms of the Node function are described in the following table.

Function Syntax	Description
<i>Node(Attribute)</i>	Returns the stored expected value of the current node. <i>Attribute</i> is an integer corresponding to an enabled payoff. This syntax can be used to report extra payoffs in Terminal Columns only.
<i>Node(Attribute; 0; LIST)</i>	Returns the expected value as above, but from a node whose path is defined by LIST. <i>Attribute</i> is an integer corresponding to an enabled payoff. In a cost-effectiveness tree, you can retrieve cost and effectiveness values as follows: Use -1 to perform CE calculations and return the new cost calculation; then use -2 to return the stored effectiveness calculation. Use -11 to return a stored cost calculation, or -12 to recalculate the node and return a new effectiveness calculation. The <i>method</i> argument 0 refers to an expected value calculation rather than a microsimulation. The <i>LIST</i> argument is a reference to the path from the root node to the node where you want calculations performed. Refer to the figure below for a description of how the path <i>LIST</i> is defined.
<i>Node(Attribute; trials; LIST)</i>	Runs a series of <i>n</i> microsimulation trials at the node whose path is defined by <i>LIST</i> . Returns the average of the trials results. The <i>Attribute</i> and <i>LIST</i> arguments function the same as for the <i>Node(Attribute; 0; LIST)</i> syntax. The <i>trials</i> argument defines the number of trials for the microsimulation. <div>  It is important to note some differences in behavior in Node() function trials, as compared to regular Monte Carlo microsimulation trials. The Node() function does not reset tracker variables to their default values before each trial (allowing trackers to be used to track information across the group of trials). By default, distributions do not resample per trial. To force distribution sampling, use tracker variables set equal to the DistForce(n) function, for example. </div>

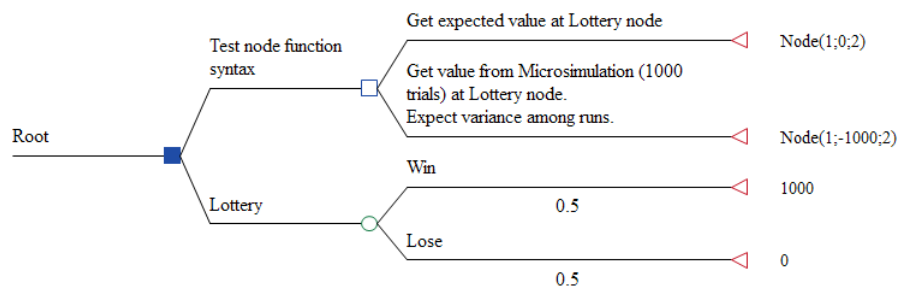
Node function syntax

The figure below includes node labels that match the appropriate *LIST* value for the Node() function to use when referring to each node.



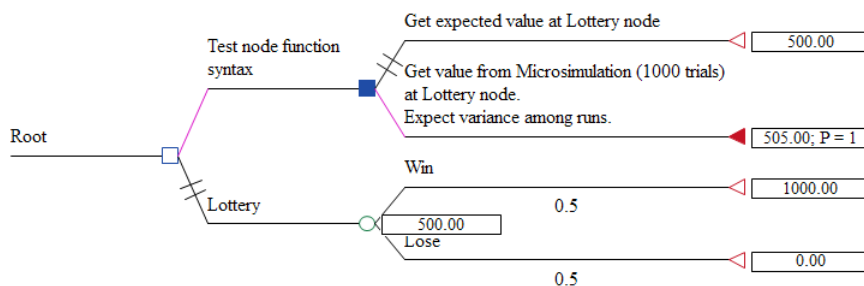
LIST values for the Node function

The Special Features tutorial example model Node Function Syntax illustrates the use of the Node function in a simple tree.



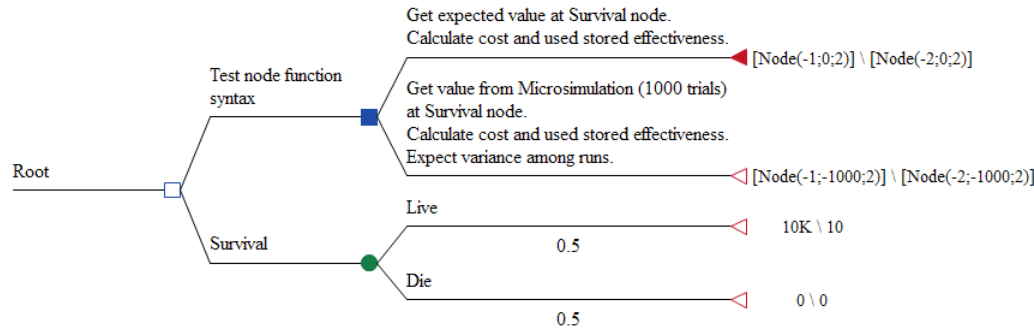
Node Function Syntax - simple model

The top two terminal nodes' payoffs use the Node function to reference the *Lottery* node via the LIST value of 2. The top payoff returns the expected value, which is always 500. The second payoff returns the value calculated by running 1000 trials at the *Lottery* node. This will return a different value near the mean of 500 each time you roll back the tree. The rolled back tree is presented below.



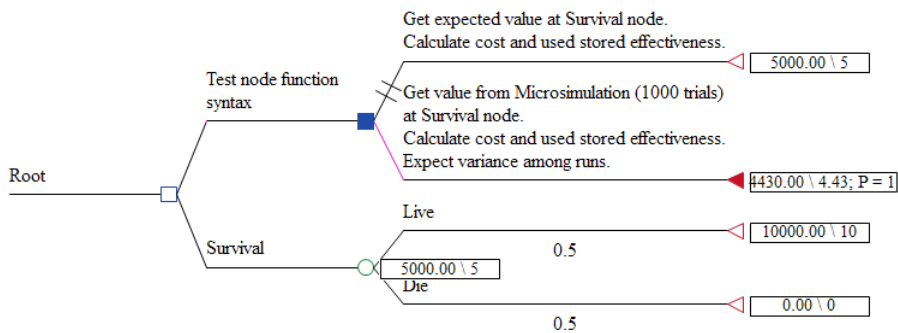
Node Function Syntax - simple model rolled back

The Special Features tutorial example model "Node Function Syntax CE" illustrates the use of the Node function in a cost-effectiveness tree.



Node Function Syntax - simple model

The top two terminal nodes' payoffs use the Node function to reference the *Survival* node via the LIST value of 2. The top cost and effectiveness payoff return the expected value for cost and effectiveness, which are always 5000 and 5. The second cost payoff returns the cost value calculated by running 1000 trials at the *Survival* node, while the effectiveness payoff returns the stored effectiveness value associated with that set of trials. This will return different cost and effectiveness values near the means each time you roll back the tree, but the cost and effectiveness values will be generated from means that are from the same set of trials. The rolled back tree is presented below.



Node Function Syntax - simple model rolled back



If the cost and effectiveness payoff sets were reversed (1 vs. 2), then you would use -11 and -12 for the attribute arguments for the Node function.

24.8 The Tree() function

The Tree() function is one of a number of TreeAge functions available which take double-quoted string arguments rather than just arguments that can be evaluated numerically. Like all functions, however, the Tree() function returns a numeric value.

The Tree() function allows you to return a calculated value from another tree. For example, you could have two separate Markov models defined for specific strategies for treating a disease. You could then have a third tree with a decision node that uses the Tree() function to refer to each strategy tree.

For the purposes of this documentation, the tree relationship is described as a *calling tree*, using the Tree function to access a *target tree*.




It is recommended that Tree() functions be carefully tested in simple models before being used in complex models.

The Tree() function syntax is described in the table below:

Function Syntax	Description
<i>Tree("target tree"; "calculation")</i>	<p>Opens a hidden copy of the specified tree or package file, and returns the results of the specified calculation.</p> <p>The target tree remains open for additional Tree() function calls.</p> <p>The "target tree" argument specifies the path and filename of the target tree to open for calculation. However, if the file is in the same directory as the calling tree, it uses just the filename.</p> <p>The "calculation" argument defines the calculation to be performed on the target tree. Usually the Node() function will be used here, in order to calculate a specific node in the target tree.</p> <p>Example usage: <i>utilXYZ = Tree("c:\my docs\projects\xyz.tre"; "Node(1;0)"</i></p> <p>The Node() function, in this case, performs a simple expected value calculation at the root node of the target tree.</p>
<i>Tree("target tree"; "calc"; 1)</i>	<p>Same as above, but if the target tree is already open and visible, the linkage is left intact after the calculation; this allows analysis commands like rollback to be used on the visible target tree for debugging purposes.</p> <p>Note that if you use this option, the variables passed to the target tree will remain in place for further calls to that tree. If you are calling the target tree with other variable definitions later in the model, do not use the third argument.</p>
<i>Tree("calculation")</i>	Can be used during an ongoing Tree() function evaluation in order to perform the given calculation at the root node of the current target tree.
<i>Tree(integer 0)</i>	Closes any open, hidden target trees. Use this, for example, if you open a target tree using the File menu and save changes to it. This will cause the next Tree() function call to reload the target tree file from disk.

Tree function syntax

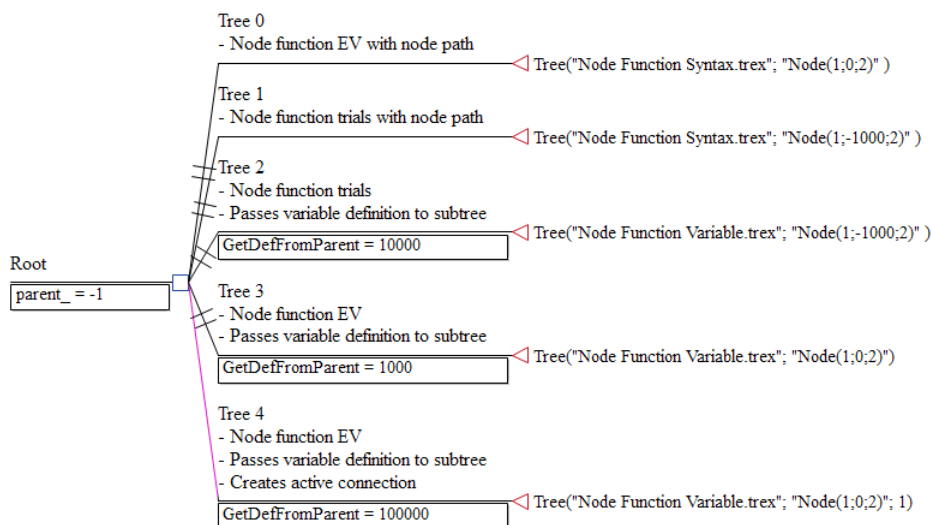
When the function opens/accesses a target tree, normally a temporary linkage is created; the target tree's root node becomes the "child" of the calling node. The power of the Tree() function is that a target tree can use variable definitions from the calling tree or to the left of the calling node. This works very much like variables in cloned subtrees. The trick is that the variable can be referenced in the target tree while having no definitions; the definitions can instead be in the calling tree. Note that the variable name must be declared in the target tree's Variables list, although it would not be defined in the target tree. The target tree must be opened before the roll back to create this active connection.

 The function's primary use is for modularization of complex trees. For example, it might offer an alternative to memory-intensive models in which hundreds or thousands of cloned copies of a Markov model are created. For example, 1000 terminal node payoffs could use the Tree() function to calculate the same Markov model, just with different variable definitions (rather than attaching 1000 clones of the Markov subtree). Calculation speed in most models will be better if the Tree() function is not used, and clones are used instead; expressions in the hidden tree that reference a variable defined in the calling tree cannot be optimized. However, using clones (or regular copies) can sometimes result in a tree whose memory requirements exceed available physical memory, resulting in an extreme decline in calculation speed. Building smaller trees linked by the Tree() function would be a good solution in this case.

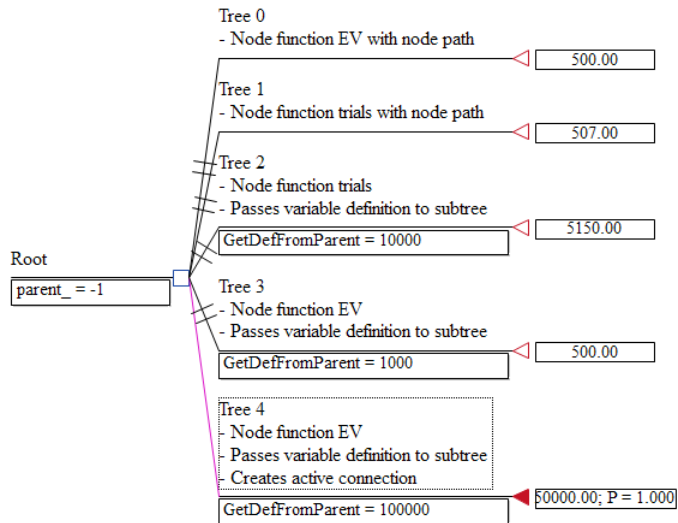
The following example uses three Special Feature tutorial examples: Tree Function Syntax, Node Function Syntax and Node Function Variable.

- Tree Function Syntax: Calling tree that uses the Tree function to call the other two trees.
- Node Function Syntax: Self-contained tree referenced at the Lottery node.
- Node Function Variable: Same as above except that it needs to receive a definition for the variable *GetDefFromParent* from the Tree Function Syntax tree.

Let's look at the Tree Function Syntax tree - both its syntax and the roll back results.



Tree Function Syntax tree



Tree Function Syntax tree - rolled back

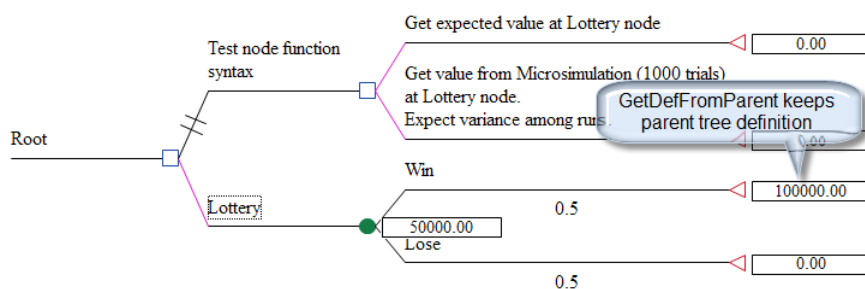
The *Tree 0* terminal node calls for the expected value of the Node Function Syntax tree at the *Lottery* node, yielding a value of 500.

The *Tree 1* terminal node runs 100 trials on the Node Function Syntax tree at the *Lottery* node, yielding a value near, but not equal to 500.

The *Tree 2* terminal node runs 100 trials on the Node Function Variable tree at the *Lottery* node. The variable definition for *GetDefFromParent* (10,000) is passed from that node to the Node Function Variable tree. This yields a value near, but not necessarily equal to 5,000.

The *Tree 3* terminal node calls for the expected value of the Node Function Variable tree at the *Lottery* node. The variable definition for *GetDefFromParent* (1,000) is passed from that node to the Node Function Variable tree. This yields a value of 500.

The *Tree 4* terminal node calls for the expected value of the Node Function Variable tree at the *Lottery* node. The variable definition for *GetDefFromParent* (100,000) is passed from that node to the Node Function Variable tree. This yields a value of 50,000. Because the Tree function contains a third argument (1), the connection from the tree to the target tree is maintained, allowing you to roll back the Node Function Variable Tree with the *GetDefFromParent* variable definition intact.



Target tree rolled back

In the figure above, the payoff expression at the Win node is *GetDefFromParent*. This evaluates to 100,000 based on the intact variable definition from the Tree Function Syntax tree.


24.9 The Command() function

The Command() function can be used to perform specialized commands.

Function syntax	Description
<code>Command("TABLES"; "table name"; "REVERSELOOKUP"; value; column)</code> <code>Command("TABLES"; "table name"; "REVERSELOOKUPTRUNCATE"; value; column)</code> <code>Command("TABLES"; "table name"; "REVERSELOOKUPCEILING"; value; column)</code>	<p>Returns index of row which matches "value" in specified column.</p> <p>If value not found, base function, truncate version and ceiling version return an interpolated index, the next lower index and the next higher index, respectively.</p>
<code>Command("GLOBALMATRIXN"; N; "GetTable"; "myTable")</code> <code>Command("GLOBALMATRIXN"; N; "GetTable"; "myTable"; startRow)</code> <code>Command("GLOBALMATRIXN"; N; "GetTable"; "myTable"; startRow; startCol)</code>	<p>Load data from a table into a Global matrix.</p> <p>N is the matrix index.</p> <p>GetTable is required for this command syntax.</p> <p>myTable is the table name.</p> <p>startRow is the matrix row to start loading data (optional).</p> <p>startCol is the matrix column to start loading data (optional).</p>
<code>Command("EXCEL"; "Open"; "Path&Workbook")</code> <code>Command("SPREADSHEET"; "Open"; Path&Workbook)</code>	Open a workbook/spreadsheet with the provided path & filename.
<code>Command("EXCEL"; "Macro"; "myMacro")</code>	Run an EXCEL macro within the opened workbook.
<code>Command("EXCEL"; "ExportGlobalMatrix")</code> <code>Command("EXCEL"; "ExportGlobalMatrixN"; index)</code>	<p>Export the non-indexed or an indexed global matrix to a new Excel workbook.</p> <p>Note that only one matrix can be exported to a new workbook. To export additional matrices, use an existing workbook.</p>
<code>Command("EXCEL"; "ExportGlobalMatrixN"; index; "Path&Workbook"; "NewSheetName"; "obsolete"; "Text_label")</code>	<p>Export Global matrix referenced by its index to an existing worksheet with the provided path & filename.</p> <p>The data will be placed in a new worksheet with the name provided.</p> <p>The sixth argument is not used.</p> <p>The Text_label argument will be presented below the data as a label.</p>
<code>Command("GLOBALMATRIXN"; 1; "ColumnLabels"; "Col1"; "Col2"; "Col3"; "Col4"; etc.)</code>	Set custom column labels for the global matrix. Usually done prior to export.

Function syntax	Description

Command function syntax

References to the workbook path must use a forward slash "/" between folders rather than a backward slash.

The Special Features tutorial example model Command Function Syntax illustrates the supported uses of the Command function. To use this example, you will need to change the path to the workbook Test Command Function.xlsm. Also, run the commands one at a time via *CONTROL+E* starting at the top.



Command Function Syntax tree

24.10 The Debug() function

The Calculation Trace Console can be very useful in debugging models. It can be used to display/report a variety of information, including: error messages, text-only analysis outputs, stored analysis summaries and detailed internal calculations.

This output is normally controlled by Tree Preferences. However, if you only want part of the overall calculations to be written to the console, use the Debug function to turn the output on and off during analysis. This is of primary interest in complex analyses when the full debugging output of calculations may be hundreds of thousands of lines (or more) per analysis.

- *Debug("1")* - Turns on detailed calculation trace.
- *Debug("0")* - Turns off detailed calculation trace.

To report only the calculations of a single probability during an analysis, for example, first turn off the debugging output of Internal calculations in the tree's Debugging preferences. Then, surround the probability by two Debug() function calls turning debugging on and then off:

See Special Features tutorial example model DebugFunction for the use of this function.

24.11 The Global() and GlobalN() functions

These advanced functions can be very powerful tools for accomplishing complex calculations and reporting tasks. It is recommended that they be tested in simple models before being used in complex models.

We recommend using the Global/GlobalN functions that are specifically designed to store, retrieve, increment, initialize and export data rather than the generic Global/GlobalN functions.

Function syntax	Description
<i>GlobalSet(row; col; value)</i> <i>GlobalNSet(matrix; row; col; value)</i> <i>Global(row; col; value)</i> <i>GlobalN(matrix; row; col; value)</i>	<p>To store and retrieve values globally, use Global() and GlobalN() to calculate "value" and set a cell in a matrix equal to the result (also returns the calculated value). Anywhere "value" is currently used in an expression, it can instead be "wrapped" in the Global function in order to save the calculated value to the matrix (for reporting purposes, for example). A value saved to a global matrix can be referenced using the second form of the Global() function; see below. The entire content of the global matrix can be dynamically saved to a text file using the third form of the function. <i>Excel Module users:</i> a simple Command("Excel";...) syntax allows you to output a matrix to a spreadsheet. <i>GlobalN() has an extra parameter matrix (integer > 0)</i> which picks a matrix to use/create, while Global() simply uses a single, default matrix (i.e., matrix 0, not matrix 1 from GlobalN). Usually a few matrices at most will be required, but thousands can be created.</p> <p>The first cell in a matrix is at row=1, column=1. Matrix size increases dynamically as cells are assigned values. Attempting to lookup a cell that is out of range, however, will result in an error. Currently, the maximum total number of cells for all matrices in the tree is 100 million (e.g., 10000x10000, 50x2000000, or 4x5000x5000). Take note of memory usage with very large numbers of matrix cells. If you are exceeding this limit, try saving/dumping a matrix to a file periodically.</p> <p><i>If both row and col are specified as 0</i>, the calculated value is instead applied to the entire existing matrix, allowing it to be dynamically reset during calculations. If an argument (including row or col) is an expression, it is evaluated at the node being calculated just as if the expression were not inside the Global() function.</p> <p>Global matrices are deleted when the tree is closed, and emptied automatically at the start of each analysis.</p>
<i>GlobalGet(row; col)</i> <i>GlobalNGet(matrix; row; col)</i> <i>Global(row; col)</i> <i>GlobalN(matrix; row; col)</i>	<p>Retrieves the value at a specified cell in a global matrix for use in a tree calculation. The first cell in the global matrix is at row=1, column=1. See information on GlobalN syntax, above.</p>
<i>GlobalIncr(row; col; incr)</i>	<p>This syntax is equivalent to: "<i>GlobalN(n;row;col; incr + GlobalN(n;row;col))</i>". If "; incr" is omitted, the increment = 1.</p>

Function syntax	Description
<i>GlobalNIncr(matrix; row; col; incr)</i>	
<i>GlobalExport(suffixNo)</i> <i>GlobalNExport(matrix; suffixNo)</i> <i>Global(value)</i> <i>GlobalN(matrix; value)</i>	<p>If <i>suffixNo</i> evaluates to a non-zero number, the matrix is silently saved to a text file (even if a simulation is running). If <i>suffixNo</i> evaluates to zero, no file is created. If <i>suffixNo</i> is less than zero, the matrix is also reported in the Debug pane (if Text-only analysis debugging setting is on). Always returns a value of 1.</p> <p>Text files are saved in the same directory as the tree, using the tree filename followed by the word "_globals_" and the integer part of <i>suffixNo</i>. A calculation in "c:\trees\mytree.tre" of the function "<i>GlobalN(3; 103)</i>" would save the current matrix #3 to a text file called: "c:\trees\mytree_globals_103.txt"</p> <p><i>Command("ExportGlobalMatrixN";n;"ColumnLabels"; "Label 1"; ...)</i> can be used to pre-set column labels.</p> <p>Excel Module users: The <i>Command("Excel";"ExportGlobalMatrixN";n)</i> function can be used to export a matrix to a spreadsheet instead. See the previous section on the <i>Command()</i> function syntax.</p>
<i>GlobalInit()</i> <i>GlobalNInit()</i> <i>GlobalInitN()</i> <i>Global()</i> <i>GlobalN(n)</i> <i>GlobalN()</i>	<p>Global matrix is emptied; size becomes 0. (Occurs automatically at the start of an analysis, or when the tree is closed. After a simulation/analysis finishes, however, matrices are still available via the above syntax.)</p> <p><i>Global()</i> and <i>GlobalN(n)</i> empty only one matrix while <i>GlobalN()</i> and <i>GlobalInitN()</i> empties all created matrices.</p>

Global/GlobalN function syntax

Most of the Global/GlobalN syntax options are used in Special Feature tutorial example models called Demo Global Functions.trex (action-specific Global functions) and Demo Global Function.trex (generic Global functions).

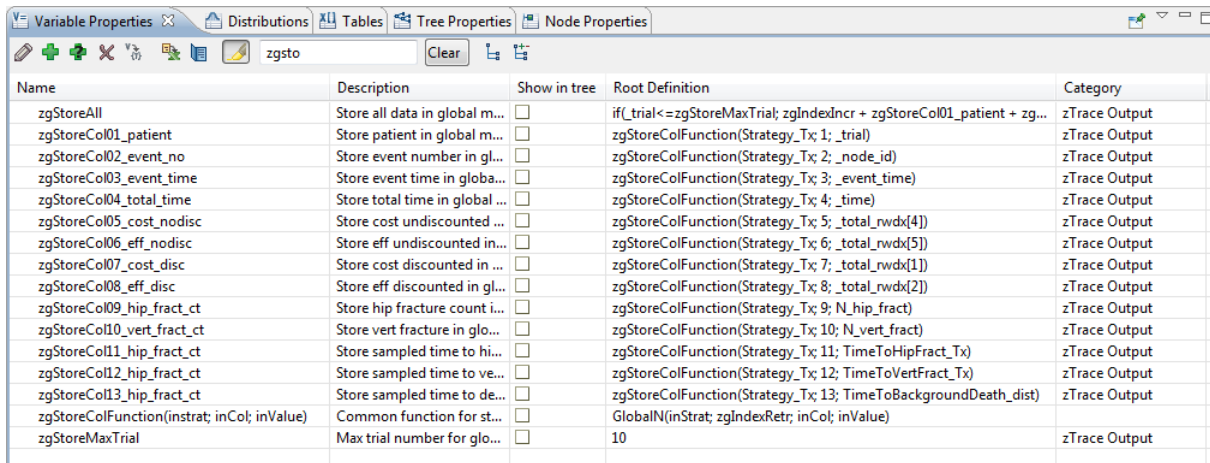


(Note: During multiprocessor simulations, Global/GlobalN refers to a set of matrices in the main thread. Separate, simultaneously processing threads evaluating a particular expression which modifies a particular matrix n will take turns updating matrix n; each thread does NOT have its own copy (except in the unusual case of a distributed simulation which divides batches among different computers.)

24.12 Custom Functions

Custom functions allow you to create a single commonly used commands/formula, which can then be used with different arguments for different contexts within the tree.

The DES tutorial example, Osteo DES Model.trex, demonstrates this technique.



Name	Description	Show in tree	Root Definition	Category
zgStoreAll	Store all data in global m...	<input type="checkbox"/>	if(_trial<=zgStoreMaxTrial; zgIndexIncr + zgStoreCol01_patient + zg...	zTrace Output
zgStoreCol01_patient	Store patient in global m...	<input type="checkbox"/>	zgStoreColFunction(Strategy_Tx; 1; _trial)	zTrace Output
zgStoreCol02_event_no	Store event number in gl...	<input type="checkbox"/>	zgStoreColFunction(Strategy_Tx; 2; _node_id)	zTrace Output
zgStoreCol03_event_time	Store event time in globa...	<input type="checkbox"/>	zgStoreColFunction(Strategy_Tx; 3; _event_time)	zTrace Output
zgStoreCol04_total_time	Store total time in global ...	<input type="checkbox"/>	zgStoreColFunction(Strategy_Tx; 4; _time)	zTrace Output
zgStoreCol05_cost_nodisc	Store cost undiscounted ...	<input type="checkbox"/>	zgStoreColFunction(Strategy_Tx; 5; _total_rwdx(4))	zTrace Output
zgStoreCol06_eff_nodisc	Store eff undiscounted in...	<input type="checkbox"/>	zgStoreColFunction(Strategy_Tx; 6; _total_rwdx(5))	zTrace Output
zgStoreCol07_cost_disc	Store cost discounted in ...	<input type="checkbox"/>	zgStoreColFunction(Strategy_Tx; 7; _total_rwdx(1))	zTrace Output
zgStoreCol08_eff_disc	Store eff discounted in gl...	<input type="checkbox"/>	zgStoreColFunction(Strategy_Tx; 8; _total_rwdx(2))	zTrace Output
zgStoreCol09_hip_fract_ct	Store hip fracture count i...	<input type="checkbox"/>	zgStoreColFunction(Strategy_Tx; 9; N_hip_fract)	zTrace Output
zgStoreCol10_vert_fract_ct	Store vert fracture in glo...	<input type="checkbox"/>	zgStoreColFunction(Strategy_Tx; 10; N_vert_fract)	zTrace Output
zgStoreCol11_hip_fract_ct	Store sampled time to hi...	<input type="checkbox"/>	zgStoreColFunction(Strategy_Tx; 11; TimeToHipFract_Tx)	zTrace Output
zgStoreCol12_hip_fract_ct	Store sampled time to ve...	<input type="checkbox"/>	zgStoreColFunction(Strategy_Tx; 12; TimeToVertFract_Tx)	zTrace Output
zgStoreCol13_hip_fract_ct	Store sampled time to de...	<input type="checkbox"/>	zgStoreColFunction(Strategy_Tx; 13; TimeToBackgroundDeath_dist)	zTrace Output
zgStoreColFunction(instrat; inCol; inValue)	Common function for st...	<input type="checkbox"/>	GlobalN(inStrat; zgIndexRetr; inCol; inValue)	
zgStoreMaxTrial	Max trial number for glo...	<input type="checkbox"/>	10	zTrace Output

Custom Functions

Note the variable `zgStoreColFunction`. The variable name itself includes a reference to arguments within parentheses, which designates this variable as a custom function. This function specifically stores data in a global matrix based on the input parameters `inMatrix`, `inCol` and `inValue`.

Note the variables `zgStoreCol01_patient`, `zgStoreCol02_event_no`, etc. Each of those variables calls the `zgStoreCol` function, with each passing the appropriate values for the input parameters.

24.13 The TrackerIncr() function

In most cases, tracker modifications in Markov simulation models are conditional. In other words, some events must occur, or one or more conditions must be met, before a tracker (i.e., a state variable) is updated. If conditional upon an event, the tracker modification ("`{T}` conditionMet=1") is put at a chance node branch. If conditional upon other trackers values, then `If()` functions or logic nodes are used within the tracker modification.

TreeAge Pro 2009 added another option: rather than creating a tracker modification at all, use the `TrackerIncr()` function in a regular variable definition.

See the note at the beginning of this section on using string concatenation to build string arguments.

TrackerIncr("tracker_name"; incr) - Adds "incr" to a current trial's value of `tracker_name`, and returns an updated value.

24.14 List function

The List function, *List(LIST)*, is used to contain a parameter list for a multivariate distribution, for example in the Dirichlet distribution.

In any tree, open the Distributions View and select a Dirichlet distribution. The list of Alphas (parameters defining the Dirichlet function) is set using the `List(LIST)` function. LIST is a set of values separated by

semi-colons used to define the Dicichlet distribution. See Distributuion Functions, Options and Types for more details about Distributions.

25. Advanced Chance Node Techniques and Options

This chapter provides information on a number of features which can be used to customize chance nodes in decision trees.

25.1 Using non-coherent probabilities

TreeAge Pro normally requires the branch probabilities of each chance node to sum to 1.0. Probabilities that meet this requirement are referred to as “coherent.” However, there are some situations where it may be useful to relax or remove this restriction (i.e., dynamic cohort analysis, parallel trials).

TreeAge Pro provides options to override the normal chance node functionality - either for the entire tree or for a specific chance node.



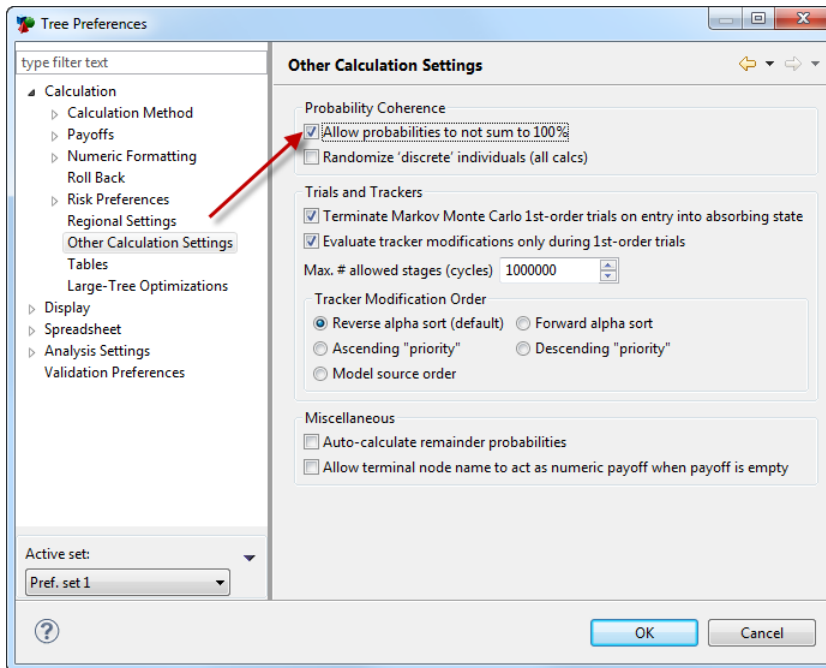
You are urged to employ these options cautiously after giving careful consideration to the potential hazards of overriding normal chance node functionality. Do not use this option simply because your probabilities are not summing to 100%. This is usually an error condition caused by probability expressions and/or distribution samples. Such an error must be corrected to generate valid analysis output.

25.1.1 Allowing non-coherent probabilities for the entire model

A Tree Preferences option allows non-coherent probabilities for the entire model. However, it is recommended that non-coherent probabilities be allowed at specific nodes instead. See next section.

To disable errors when using non-coherent probabilities:

- Choose Tree > Tree Preferences from the menu or click *F11* to open the Tree Preferences Dialog.
- Navigate to the preference category Calculation > Other Calculation Settings.
- Check the option "Allow probabilities to not sum to 100%".



Tree Preferences - Allow Non-coherent Probabilities

As precautions against unintended use, the status bar at the bottom of the TreeAge Pro window displays text to notify you that this setting is active.

CALC: Payoff 1, Non-Coherent Probs

Status bar - non-coherent probabilities



Non-coherent probabilities are not compatible with microsimulation (first-order trials) unless integer values are used for the probabilities for parallel trials, but can be used in probabilistic sensitivity analysis using expected value calculations.

25.1.2 Allowing non-coherent probabilities at a specific node

You can also allow non-coherent probabilities at specific chance and Markov nodes in the model; other node types do not allow this option. This is a better approach since TreeAge Pro will still reject non-coherent probabilities at other nodes in the model.

To change the probability coherence option for a node via the context menu:

- Right-click on a chance node or a Markov node.
- Within the context menu, choose Change Coherence > your desired option.

To change the probability coherence option for a node via the main menu:

- Select a chance node or a Markov node.
- Choose Node > Probability Coherence > your desired option.

The coherence options available are:

- *Use Tree Preference setting*: Do not override for this node. Use the option from tree preferences.
- *Allow < > 100% for this node*: Allow non-coherent probabilities at this node.
- *Do not allow < > 100% for this node*: Do not allow non-coherent probabilities at this node.
- *Normalize probabilities*: Normalize probabilities to force them to sum to 100%.

Probability coherence options

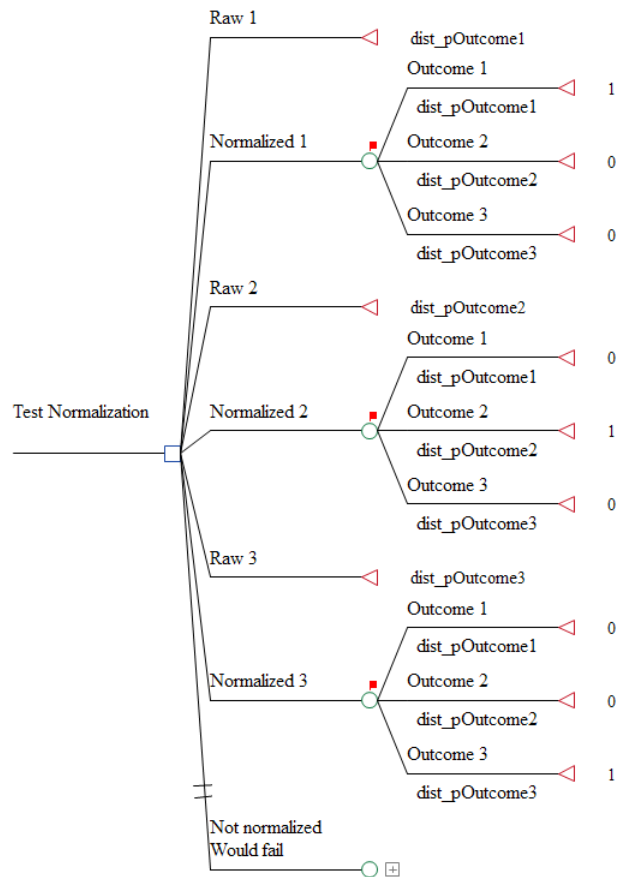
If the probabilities are allowed to be non-coherent, you will get odd expected values. This is not recommended, except when you need to add/subtract from the cohort size (dynamic cohort modeling).

Normalization will take the probabilities that do not sum to 100% and normalize them to make them sum to 100%. For example, if you had probabilities 0.3, 0.4 and 0.5, the total sum would be 1.2. By dividing each probability by 1.2, the probabilities would be normalized.

	Original	Normalized
Prob1	0.3	0.25
Prob2	0.4	0.333333
Prob3	0.5	0.416667
Sum	1.2	1.0

Prob normalization

The Special Features tutorial example model ProbabilityNormalization demonstrates this technique.



Probability normalization tree

The first 6 strategies alternate between probabilities sampled from a distribution and the normalized probabilities. The *Raw 1* strategy simply reports the first probability sampled from a distribution. The *Normalized 1* strategy returns the probability after it has been normalized at the chance node.

The bottom strategy is excluded from the analysis because it would fail since the probabilities are not normalized.

Note the red flags above the nodes with probability coherence overrides. This indicator helps identify when a node will function differently from the rest of the model. Node branch comments are automatically created with the text "(Node Probability Override)", which in turn causes the flags to be presented.

Run Monte Carlo Simulation/Sampling on the model to see the original and normalized probabilities. The summary mean values below illustrate the normalization.

Statistic	Raw 1	Normalized 1	Raw 2	Normalized 2	Raw 3	Normalized 3
Mean	0.54	0.52	0.21	0.20	0.30	0.28
Std Deviation	0.15	0.11	0.11	0.09	0.13	0.10
Minimum	0.21	0.27	0.02	0.02	0.08	0.11
2.5%	0.24	0.31	0.03	0.03	0.09	0.12
10%	0.36	0.37	0.08	0.09	0.13	0.14
Median	0.55	0.53	0.20	0.19	0.29	0.27
90%	0.72	0.67	0.36	0.32	0.49	0.41
97.5%	0.78	0.72	0.44	0.41	0.61	0.49
Maximum	0.80	0.74	0.61	0.44	0.64	0.57
Size (n)	100.00	100.00	100.00	100.00	100.00	100.00
Variance	0.02	0.01	0.01	0.01	0.02	0.01
Variance/Size	0.00	0.00	0.00	0.00	0.00	0.00
SQRT[Variance/Size]	0.01	0.01	0.01	0.01	0.01	0.01

Normalization tree analysis results

Note that the original mean value for probabilities 1, 2 and 3 are 0.54, 0.21 and 0.3, while the normalized mean values are 0.52, 0.20 and 0.28.

Contrast this with results from the Special Features tutorial example model ProbabilityNonCoherence. In that model, no normalization is done; the bad probabilities are simply allowed. This is not recommended.

25.1.3 Reporting future net present values in the tree

Discounting is normally used to generate the net present value of future values. In nearly all cases, the net present value of future values is less than the future value itself due to inflation. Decision analysis is then performed using the net present values for all values in the model.

If a decision is to be made at an intermediate node in the tree which occurs in the future (e.g., options valuation), it may be necessary to calculate the net value at the time of the future decision rather than the present time. This would require discounting from the distant future value back to the less distant decision point rather than back to the present.

In situations where it is appropriate, it is possible to use non-coherent probabilities to perform the discounting instead of discounting each terminal node payoff. Net present value discounting is generally performed by dividing by $(1 + \text{discount_rate})^{\text{time}}$. By turning off probability error checking, this division could instead be performed on each branch probability. If done correctly, the expected values calculated at the root node of the tree will be unchanged from a standard, payoff discounting model. See the TreeAge web site for more information.

As with other uses of probability non-coherence, care must be taken to ensure that hidden probability errors are not unknowingly introduced.

25.1.4 Discrete and dynamically-sized Markov cohorts

Normally, decision trees are used to calculate an expected, or average, value. In budget-oriented or population-based modeling, however, the ultimate goal may be to determine not an average, but an overall cost or benefit. In some cases, this can be accomplished simply by multiplying the expected

value by some number (e.g., the number of projects, or the size of a population). If TreeAge Pro's probability coherence requirement is turned off, an equivalent option would be to do the multiplication via tree "probabilities."

Non-coherent probabilities might be used to model a population whose size changes over time. For example, in a Markov model built using the Healthcare module, the starting population can be initialized — sized and distributed among possible health states — prior to the Markov calculation. Then, during Markov calculations, non-coherent probabilities could be used to change the size of the population, modeling entry from other populations (i.e., from uninfected to infected) or internal population growth (births).

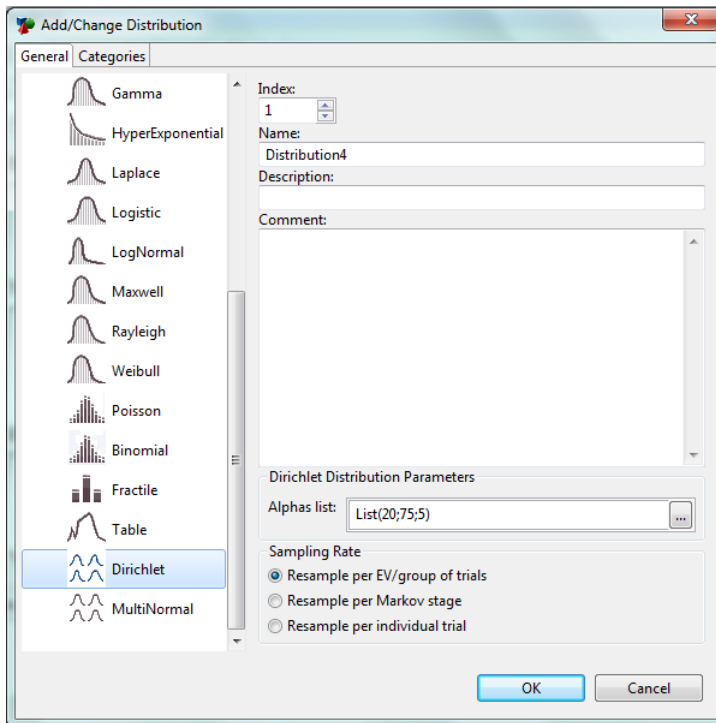
When allowing non-coherent probabilities, a sub-option to randomize "discrete" individuals is available that will maintain integer probabilities at subsequent chance nodes — in effect, keeping individuals whole by randomizing them at chance nodes (during any analysis, not just simulation). This might be relevant, for example, where small probability events are critical (i.e., in a vaccination model where continued transmission of a contagious disease requires a "whole" carrier).

25.2 Sampling probabilities from a Dirichlet distribution

If a chance node has more than two branches, performing a sensitivity analysis or Monte Carlo simulation that changes the values of these probabilities can be problematic.

One option is to write expressions that normalize the chance node's probability expressions. For example, let's say the node has three outcomes: A, B, and C. Rather than assigning variables to two probabilities, and using the # remainder calculation in the third, you could do the following: assign three expressions that always sum to 1.0, like $pA/(pA+pB+pC)$ and $pB/(pA+pB+pC)$ and $pC/(pA+pB+pC)$. No matter what values (≥ 0) are assigned to pA, pB, and pC, the three normalized probabilities will always sum to 1.0. (The # remainder could still be used in place of one of these.)

TreeAge Pro offers another solution using a special multivariate form of the beta probability distribution, the Dirichlet distribution. This distribution can be used to represent the uncertainty in all of the probabilities of a chance event. During Monte Carlo simulation, the distribution can sample probabilities for each branch, while ensuring that probabilities sum to 1.0.



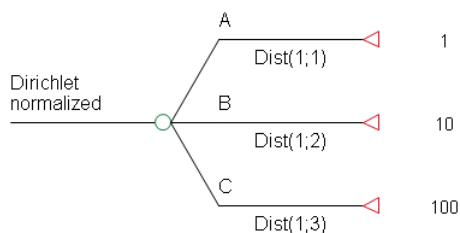
Dirichlet distribution

If the distribution is parameterized with a list of N *alpha* values, N independent $\text{Gamma}[\text{alpha}_n, \text{beta}=1]$ distributions will be sampled and the samples normalized to create a list of N probabilities which are guaranteed to sum to 1.0.



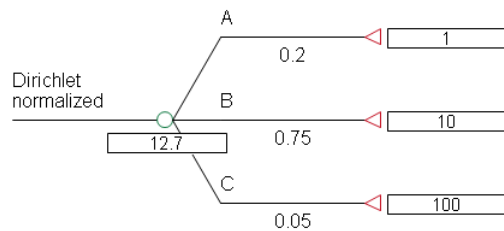
Note that using a larger total sum of the alpha values will reduce the variation of each probability from its mean. For example, $\text{List}(200; 750; 50)$ generates the same three mean values as $\text{List}(20; 75; 5)$, but the sample sets will tend to be closer to the mean values. Note that you can use a Dirichlet distribution without sampling to normalize probabilities for a deterministic sensitivity analysis (not PSA). The mean values for each value generated from a Dirichlet distribution is equal to the individual List argument divided by the sum of all the List arguments. Therefore, if you adjust one of the List values for deterministic sensitivity analysis, the other values will adjust proportionally to keep the overall probabilities coherent.

To utilize the sampled probabilities in the model, the Dist function is used with a second argument indicating which branch (i.e., alpha) to use, as shown below.



Dirichlet tree

Rolling back the tree shows the mean values of the probabilities, which are simply the normalized alpha parameters.



Dirichlet tree rolled back

Performing a simulation in the example tree shows the effect of sampling independent Gamma distribution values, based on the list of alpha parameters, and then normalizing. For each iteration of the simulation, a different set of Gamma random variates is drawn. Each iteration results in a different sum, as well as different ratios of the Gamma random variates to the sum (i.e., the probabilities). Normalization ensures that the resulting probabilities sum to 1.0.

Running PSA on the model yielded the following results.

Statistic	Dirichlet normalized
Mean	12.507
Std Deviation	1.874
Minimum	9.15
2.5%	9.337
10%	10.085
Median	12.154
90%	15.125
97.5%	16.286
Maximum	16.989
Size (n)	100
Variance	3.512
Variance of Mean (var/n)	0.035
Std Error of Mean	0.187

Dirichlet tree PSA

Note that the mean value from the 100 iterations is close to the roll back expected value. However, the variance among the iterations reflects the different probabilities that are generated by the Dirichlet distribution. The Values, Dists, Trackers link shows the three individual probabilities generated from the distribution for each iteration. Note that the first, second and third values from the distributions are near 0.2, 0.75 and 0.05 respectively as expected.

Monte Carlo Simulation Report				
ITERATION	EV	DIST_1_VAR_1	DIST_1_VAR_2	DIST_1_VAR_3
1	14.018	0.203	0.732	0.065
2	13.279	0.22	0.722	0.058
3	11.931	0.255	0.698	0.047
4	13.385	0.167	0.779	0.054
5	11.317	0.2	0.765	0.035
6	11.188	0.249	0.713	0.038
7	13.348	0.199	0.744	0.057
8	16.004	0.238	0.672	0.09
9	13.948	0.224	0.71	0.066
10	10.44	0.222	0.751	0.027
11	13.452	0.175	0.77	0.056
12	11.67	0.113	0.857	0.03
13	13.607	0.126	0.821	0.053
14	13.178	0.248	0.692	0.06
15	14.985	0.228	0.694	0.078
16	16.443	0.179	0.732	0.089
17	10.784	0.182	0.791	0.027
18	14.774	0.186	0.742	0.072
19	13.586	0.214	0.725	0.061
20	10.853	0.207	0.763	0.03

Dirichlet tree Values, Dists, Trackers output



In TreeAge Pro, simulation text reports and graphs only report the first sampled Dirichlet probability. However, it is possible to use either a tracker variable or TreeAge Pro's Global() matrix function to store and report each sampled probability. For example, to create a table of a branch's sample values, wrap the Dist() function reference in the Global() function, as in:
Global(_sample; branch; Dist(1; branch))

26. Bayes' Revision in Decision Trees

This chapter shows how, in the tree window, TreeAge Pro can assist you in performing the calculations that implement probability revision using Bayes' theorem.

26.1 An introduction to Bayes' revision

If your model includes imperfect tests or forecasts followed by decisions, you may wish to utilize TreeAge Pro's Bayes' revision feature. TreeAge Pro can automatically perform probability revisions using Bayes' theorem. The process occurs once during the initial construction of the model; based on your answers to a few questions, TreeAge Pro will generate a set of variable definitions that calculate the revised probabilities. The probability expressions will be recalculated every time the model is evaluated and results can change as your estimates of prior and likelihood probabilities (see below) change.

Bayes' revision is implemented in both the tree and influence diagram windows. Bayes' revision in the tree window is able to revise probabilities automatically based upon a single test. To revise probabilities associated with sequential tests, you should initially build your model as an influence diagram.

There are two methods of applying Bayes' theorem on your model, each of which is described later in this chapter:

1. For test results using sensitivity and specificity.
2. For a general m-by-n grid.

26.1.1 Probability revision using Bayes' theorem

Bayes' revision allows decision makers to calculate *decision* probabilities from *likelihood* probabilities. Likelihood probabilities, or forecast likelihoods, are answers to questions such as, "If this test is performed on a part *known* to be faulty, what is the probability of a positive result indicating a problem?" This type of probabilistic information is often available, but is not immediately useful in making decisions. What's needed are the decision probabilities, which address questions such as, "If a particular part tests positive, what is the probability that is really faulty?"

The decision probabilities are so named because, in the real world, they are the probabilities upon which decisions are based. These are also sometimes called posterior (or *a posteriori*) probabilities.

The basic formula for revising probabilities is:

$$P(H | E) = \frac{P(E | H) * P(H)}{P(E)}$$

where H is the hypothesis (e.g., faulty or not faulty) and E is the evidence (e.g., test result). The formula is applied once for each hypothesis-evidence combination — for example, $P(\text{not faulty} | \text{positive})$, or the

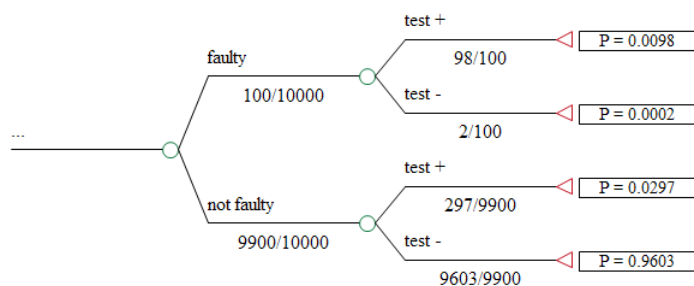
probability not faulty given a positive test result. $P(H)$ represents the prior (or *a priori*) probability of the condition. $P(E)$ is a *marginal* probability, calculated as part of the revision.

26.1.2 A simple numeric illustration

The following example is designed to offer a sense of the potential usefulness of Bayes' revision. If you are already familiar with the types of applications that require Bayes' revision, you may want to skip this section.

Consider an automated test for a defect in a semiconductor. The defect is present in 1% of the items under scrutiny. It has been demonstrated that the available test will detect 98% of the faulty materials, meaning that 2% of those pieces with the defect will not be picked up by the test. Also, the test is known to incorrectly identify as faulty 3% of those pieces that are without defect.

You have considered installing a machine to perform this test in your facility. What is the likelihood that a part which tests positive actually has the defect? How certain can you be of parts that tested negative don't have a defect? The information about the accuracy of the testing equipment provided above does not directly answer these crucial questions.



Bayes' Revision illustration - part 1

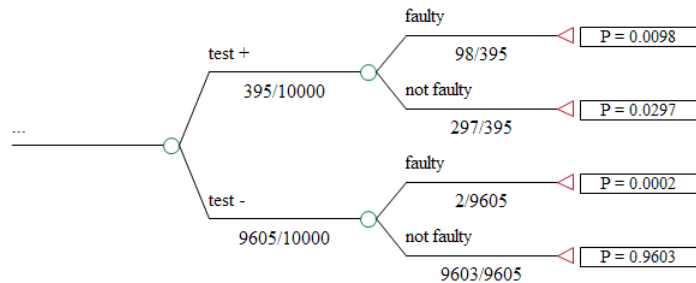
Let's say we have a batch of 10,000 items to be tested.

If the estimated prior probability of defect is 1%, we would expect 100 items in the batch to have the defect. Of these, about 98 should test positive. Of the 9,900 pieces without the defect, we said approximately 3% (297) would test positive.

Thus, a total of 395 (297 + 98) test subjects would test positive (this is one of the marginal test probability).

The Bayes' revision formula is intuitive when illustrated and worked out using a tree, as shown below.

The first revised decision probability is the ratio 98/395, or approximately 25%. This is the probability that a positive test actually indicates the presence of the defect. In this case, 75% of the positive tests are in error. The other decision probabilities are similarly calculated. With this information, a decision maker could compare the performance of the new test with existing methods or competing technologies.



Bayes' Revision illustration - part 2

26.2 Bayes' Revision - Sensitivity/Specificity

The most common form of Bayes' revision performed in healthcare models relates to imperfect tests using sensitivity and specificity. In a typical model, you screen for a condition, but the test yields some false positives and false negatives. Reported data on the sensitivity and specificity of a test and the prevalence of the disease can be integrated into a model as parameters, which are revised to generate dependent probabilities required for the model.



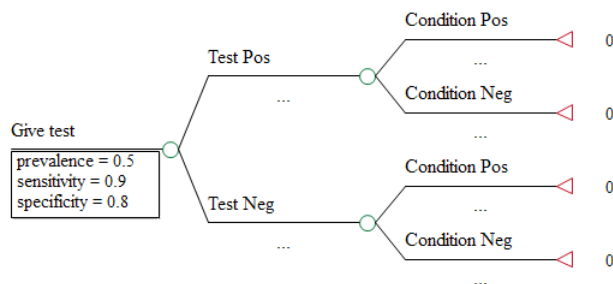
Sensitivity and specificity are statistical measures of the performance of a binary classification test.

Sensitivity (also called the true positive rate) measures the proportion of actual positives which are correctly identified as such (e.g. the percentage of sick people who are correctly identified as having the condition).

Specificity (also called the true negative rate) measures the proportion of negatives which are correctly identified as such (e.g. the percentage of healthy people who are correctly identified as not having the condition).

For this version of the Bayes' revision, you must first create the model structure to mirror the test results and actual condition. This structure can be anywhere in the tree, but it must follow the structure of a chance node with two branches that are each also chance nodes with two branches.

The Health Care tutorial example model, BayesSensSpec-start.trex, is presented below.



Bayes' sensitivity/specificity example model - before revision

The model contains the standard test and condition node structure as well as the three variables prevalence, sensitivity and specificity.

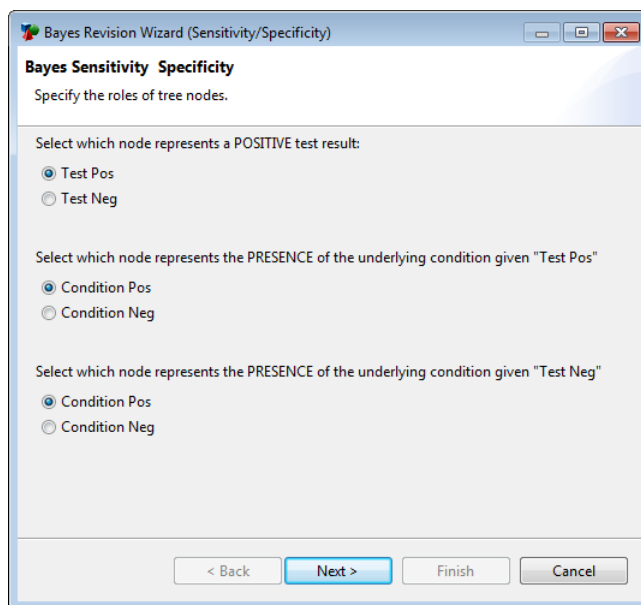
To initiate the Bayes' revision using sensitivity and specificity:

- Select the Give test chance node.
- Choose Subtree > Bayes' Revision > Sensitivity, Specificity from the menu.

You are then presented with a wizard dialog to walk you through the revision process. In the first step, you simply select the test positive node. Then select the two condition positive nodes that follow the positive and negative test nodes. The dialog uses the node label text to guide you through the process. Then click Next.



The radio button for the top branch is always selected in the dialog below. Do not assume that this is the correct selection for your model.



Bayes' sensitivity/specificity wizard - select nodes

Then you select variables to for the prevalence, sensitivity and specificity values required for the revision. The wizard includes a "Show variable definitions" link if you need to create/define these variables on the fly. In our example, these variables were already defined, so they can simply be selected.



Note that if you create variables on the fly through the mini Variable Definitions View within the Wizard, definitions will be placed at the "root" of the revision, which may not be where you want them. You might choose to create the three required variables first, so you can simply reference them within the wizard.

The wizard also contains a "Show Bayes grid" link to allow you to see the calculated values that will be used in the model.

Bayes Revision Wizard (Sensitivity/Specificity)

Specify Sensitivity, Specificity and condition probability.

Prior probability of "Condition Pos": prevalence

SENSITIVITY of the test: sensitivity

SPECIFICITY of the test: specificity

[Show Variable Definitions](#) (for reference) [Hide Bayes Grid](#)

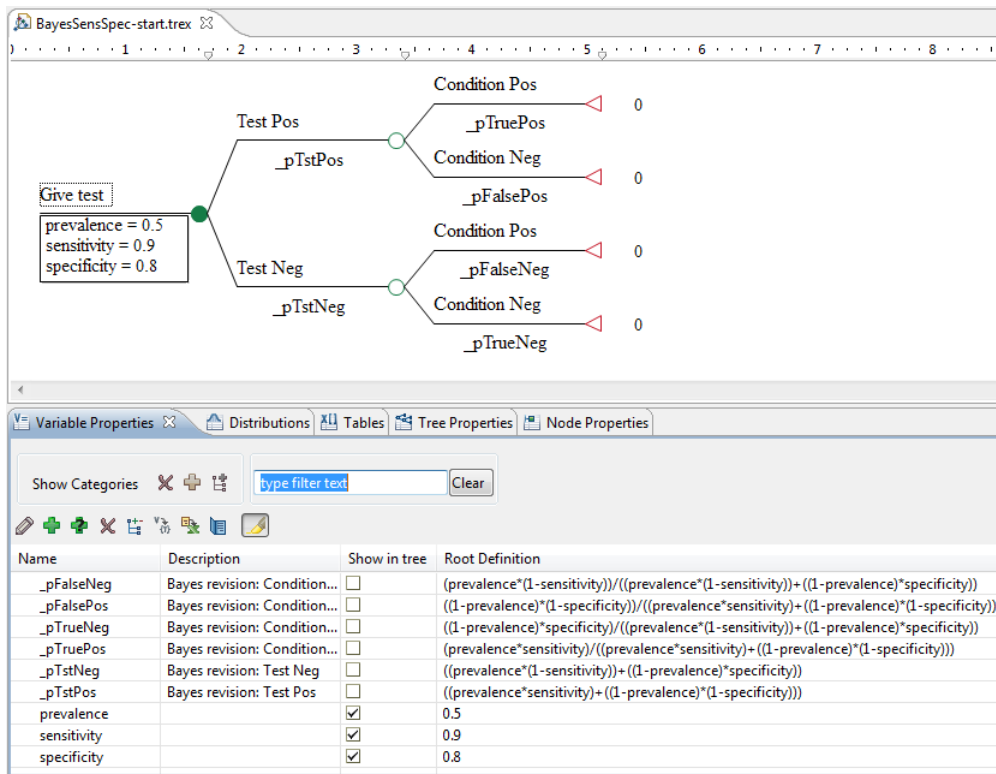
		Test Pos	Test Neg
Marginal Probabilities:	Test:	0.55	0.45
Inverted Conditional Probabilities:	Condition Pos	0.818181818	0.111111111
	Condition Neg	0.181818182	0.888888889
Joint Probabilities:	Condition Pos	0.45	0.05
	Condition Neg	0.1	0.4

[Refresh Bayes Grid](#)

< Back Next > Finish Cancel

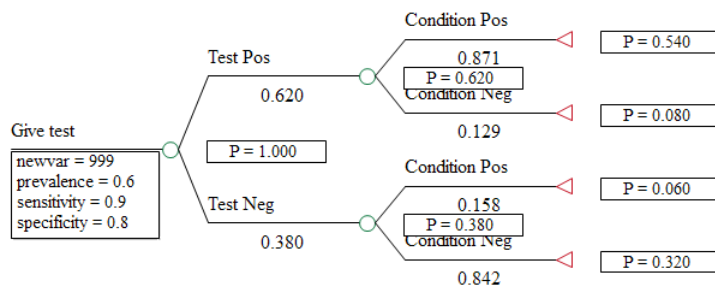
Bayes' sensitivity/specificity wizard - select probability variables

When you click Finish, the revised probability variables are then defined at the "root" revision node with references to the three independent variables. The revised variables are then placed into the tree's branch probabilities. The revision variables are set to be hidden in the tree structure.



Tree after Bayes' revision with sensitivity, specificity

The Health Care tutorial example model BayesSensSpec-end.trex includes the final set of revised probabilities. Roll back shows all the calculated revised probabilities in the model.



Revised model rolled back

26.3 Bayes' Revision - Grid

The grid approach to Bayes' revision also revises the known probabilities to create the probabilities needed for the model. However, it differs in two significant ways:

1. It supports any m-by-n sized grid.
2. It generates the tree structure rather than referring to an existing tree structure.

To use the grid Bayes' revision, you should first obtain numeric values for the likelihood probabilities associated with the test and the *a priori* probabilities for the hypotheses. Then, use the grid revision wizard to create the model's structure and probabilities.

This chapter will examine the contents of an oil well and a test for seismic soundings. We will assume that we have knowledge about wells in a target area based on prior success in the immediate region. Based on that knowledge, we estimate the likelihood that the well has no oil, some oil or lots of oil as shown below.

Condition	Probability
No Oil	5/10 (50%)
Some Oil	3/10 (30%)
Lots of Oil	2/10 (20%)

Based on prior experience, a table is constructed showing what we can expect for test results (test neg, test pos moderate, or test pos high) based on a well's actual state (no oil, some oil, or lots of oil) as shown below.

Condition	Test Low	Test Med	Test High	Total
Given No Oil	0.6	0.3	0.1	1.0
Given Some Oil	0.3	0.4	0.3	1.0
Given Lots of Oil	0.1	0.4	0.5	1.0



This example uses the default 3x3 grid. Please note that the grid does not need to be symmetrical.

The starting point for your Bayes' revision is a chance node with no branches. The subtree generated by the process can then be copied or moved later as needed. The Oil tutorial examples model, BayesGrid-start.trex, is a quick starting point for working with the grid Bayes' revision.

To start the grid Bayes' revision:

- Open the BayesGrid-start.trex model.
- Select a chance node with no branches.
- Choose Subtree > Bayes' Revision > M by N grid from the menu.

The Wizard then opens.

Bayes' Revision Dialog

Bayes' Revision
 ⓘ Fill all the enabled empty field values of the independent probabilities

Conditions number Results number

Condition Description	Condition Short Name	Variable Definition	Test Result Short Name	Result 1	Result 2	Result 3
Condition 1	Cond1	0	Given Condition 1	0	0	1.0
Condition 2	Cond2	0	Given Condition 2	0	0	1.0
Condition 3	Cond3	1.0	Given Condition 3	0	0	1.0

[Show Bayes Grid](#)

Bayes' revision grid wizard

Note the two spinners at the top that can be used to change the shape of the grid.

The wizard provides entry fields to the left for the condition descriptions (for node labels), for condition short names (for variable names) and variable definitions (for values). It also provides entry fields for test results (for node labels), for test result short names (for variable names) and given conditions (for values).

Based on the information provided earlier, the grid could be filled out as seen below. The "Show Bayes Grid" link has been clicked to show all the Bayes' revision calculated values.

Bayes' Revision
 ⓘ Fill all the enabled empty field values of the independent probabilities

Conditions number Results number

Condition Description	Condition Short Name	Variable Definition	Test Result	Test Low	Test Med	Test High
			Test Result Short Name	TestLow	TestMed	TestHigh
No Oil	NoOil	0.5	Given No Oil	0.6	0.3	0.1
Some Oil	SomeOil	0.3	Given Some Oil	0.3	0.4	0.3
Lots of Oil	LotsOil	0.2	Given Lots of Oil	0.1	0.4	0.5

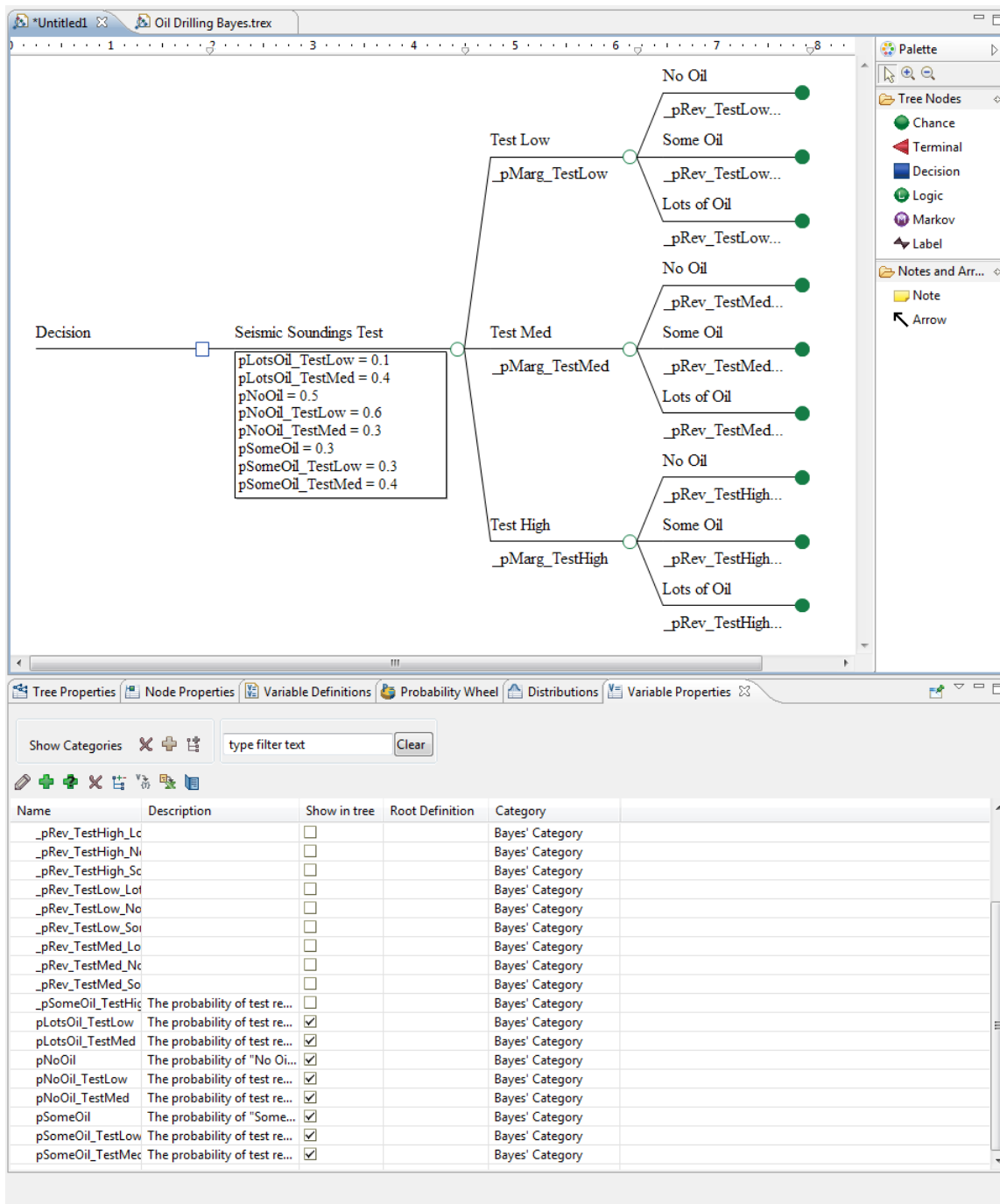
[Hide Bayes Grid](#)

		Test Low	Test Med	Test High
Marginal Probabilities:	Test:	0.41	0.35	0.24
Inverted Conditional Probabilities:	No Oil	0.732	0.429	0.208
	Some Oil	0.22	0.343	0.375
	Lots of Oil	0.049	0.229	0.417
Joint Probabilities:	No Oil	0.3	0.15	0.05
	Some Oil	0.09	0.12	0.09
	Lots of Oil	0.02	0.08	0.1

[Refresh Bayes Grid](#)

Bayes' revision grid wizard with values

Then click Create Bayes Subtree to create the subtree with all independent, conditional and calculated variables created and defined.



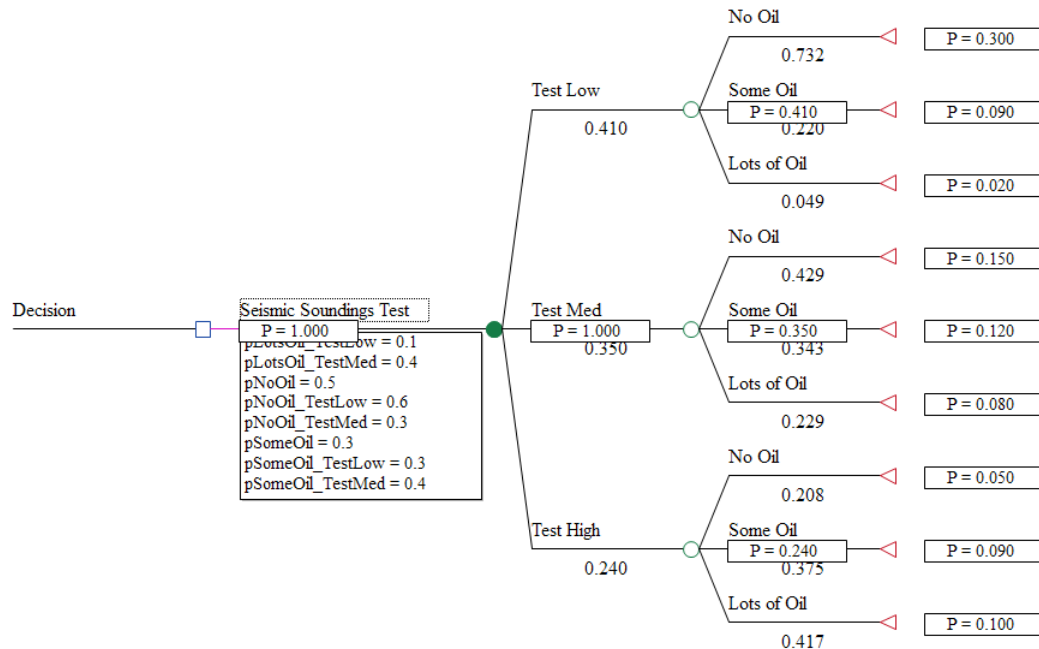
BayesGrid-end.trex

Note the following elements of the subtree that is created.

- The tree structure is built automatically based on the M by N grid.
- The node labels are populated based on the Condition Description and Test Result labels entered in the grid.
- The independent variables are named based the short names entered for the condition and test results and are defined based on values in the grid.

- The revised probability variables are named starting with an underscore and are not "shown in tree." They are defined using Bayes' revision theorem.

The Oil tutorial example, BayesGrid-end.trex, is a tree after the revision with the end nodes changed to terminal nodes to display the path probabilities, as shown below while rolled back.



BayesGrid-end rolled back

Note that the path probabilities, conditional probabilities and test result probabilities match the values presented in the grid.



The names and values entered in the last Bayes grid are stored with the model. This allows you to create another Bayes subtree using the same data. The Oil tutorial examples model BayesGrid-start-populated.trex is pre-populated with the Bayes grid data described above.

27. Utility Functions and Risk Preferences

TreeAge Pro allows you to set up a risk preference utility function in a tree, which can be used to account for a decision maker's aversion to risk.

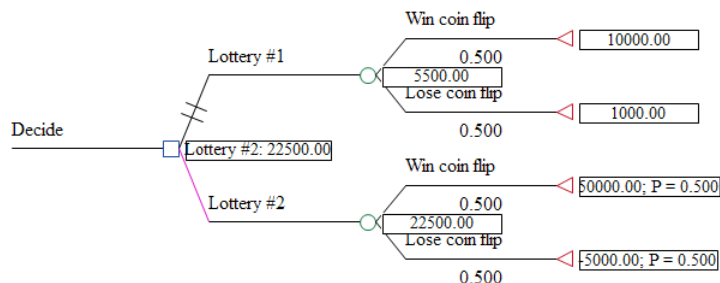
27.1 Risk preference: an illustration

Most decision makers are risk-averse to some degree. They are willing to pay a premium, small or large, to avoid risk. The decision maker's risk preference can be incorporated into a decision tree.

Assume that a rich uncle offers you an opportunity to win some money. He proposes to flip a coin giving you the opportunity to receive either \$10,000 or \$1,000, depending on whether you correctly predict the outcome. If you call the flip correctly, you will receive \$10,000, and if you are wrong you will receive \$1,000.

To make this game more interesting, assume that your uncle complicates matters by offering an alternative opportunity. The alternative is also a coin flip. Under this one, you will receive \$50,000 if you are correct, but you will have to pay him \$5,000 in the event you lose on the coin flip. There will be only a single coin flip; it is up to you to choose between the two. As you will see, it may not be wise to base your decision solely on traditional expected value calculations.

The tree in Business tutorial examples, Risk Illustration, models your uncle's offer.



Risk Illustration tree

As the tree illustrates, there are two lotteries. Both provide the same (50 - 50) odds of winning, but they have different outcomes. You must choose one of them. On the basis of expected value, you should choose lottery #2. Its expected value (\$22,500) is more than four times that of lottery # 1 (\$5,500).

However, what about the risk posed in lottery #2 that you could actually end up losing \$5,000? At least in lottery #1 there is no risk of being out-of pocket – you are guaranteed to win something. How one responds to the downside risk posed by lottery #2 involves a subjective analysis of the decision maker's *aversion to risk*.

27.1.1 Certainty equivalents and risk aversion

Consider lottery #1 described above. The expected value is \$5,500. Would you sell the opportunity to play this lottery for \$4,000? If you were offered \$3,000 by a third party who wanted to buy into the lottery, would you sell?

The minimum value for which you would sell the lottery is your certainty equivalent for this lottery. The certainty equivalent of a lottery can be perceived as the expected value of that lottery, adjusted for risk preference (the risk-adjusted expected value).

A certainty equivalent is similar to an expected value, in that it is a single numeric quantity which represents the value of an uncertain event. The certainty equivalent is a subjective measure; it is the answer to the question, "What is the minimum (or maximum) value for which I would trade this uncertainty?"

Now consider a situation which is undesirable from the start. Lottery #3 is a coin flip in which you will either owe your uncle \$2,000 or you will owe him \$12,000. In this situation, we are interested in finding the maximum amount that you are willing to pay to a third party to assume your obligation under the lottery. Would you pay \$4,000? Or \$5,000? Your answer to this question is your certainty equivalent for that lottery.

The certainty equivalent for a lottery is usually in the same numeric range as the expected value. The gap between the certainty equivalent and the expected value is a measure of risk aversion.

Most decision makers are risk-averse to some degree. They are willing to pay a premium, small or large, to avoid risk. Their certainty equivalent for any lottery will be lower than the lottery's expected value. In contrast, a risk-seeking decision maker is one whose certainty equivalent for a lottery is higher than the lottery's expected value. The risk taker is willing to pay a premium in order to participate in the lottery.


27.2 Risk preference curves

A straight-line risk-preference curve represents a decision maker who is risk-neutral. This type of decision maker bases decisions on expected values rather than certainty equivalents.

A risk-averse decision maker will have a curve with a decreasing slope, meaning that certainty equivalent is less than expected value. The curve will typically be steeper in the low value range, where aversion to risk is weak, and will grow progressively flatter as the values get larger (both positive and negative), where aversion to risk becomes stronger. The more risk-averse you are, the more your curve will deviate from the 45° straight line representing risk neutrality.

If you encode a curve that includes some unexpected bumps when graphed, this means that some of your responses were inconsistent. You should repeat the process. Don't be discouraged; developing a meaningful non-constant risk utility curve takes hard thinking and careful consideration.

For further reading on risk preference, refer to the general texts on decision analysis listed at the end of Chapter 1.

 Risk preference functions can be used only if the calculation method is set to Simple.

27.3 Creating a risk preference function

TreeAge Pro is able to record your risk function as a mathematical curve and apply this curve to the expected value of an uncertainty. Recommendations are then made based on your derived certainty equivalents, rather than on expected values.

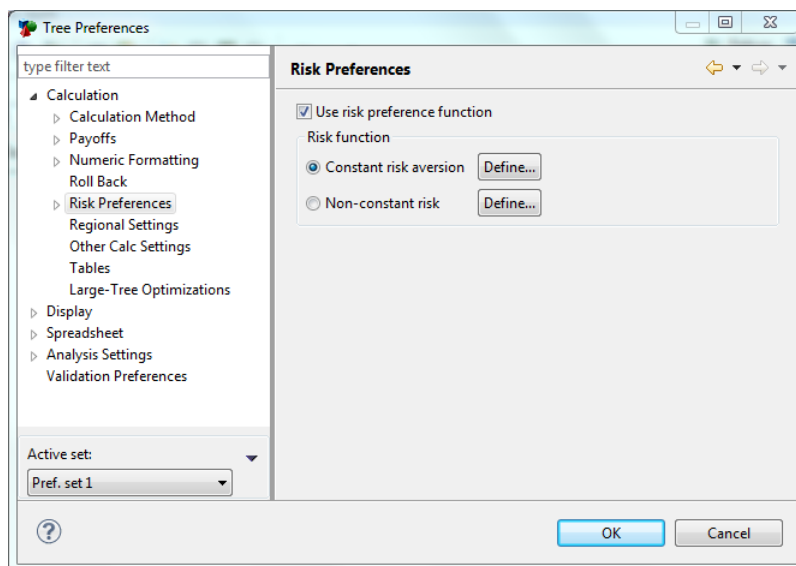
There are two types of curves, or risk functions, which TreeAge Pro can use. The *constant risk aversion* function is calculated using the formula:

$$U(x) = 1 - e^{(-x/r)}$$

where U is an arbitrary utility scale, and R is a risk preference coefficient, described below. The utility scale is used only for internal calculations; the formula's inverse is later applied to find certainty equivalents.

The *non-constant risk function* is tailored to fit your specific model, and so is superior to the constant risk aversion function in many respects, except that it takes a little longer to set up initially. TreeAge Pro will ask you a series of questions about your certainty equivalents for the model you are working on. It will then create a curve made up of line segments approximating your true risk function.

Both types of risk functions are entered via Tree Preferences.



Tree Preferences - Risk Preferences

To use risk preferences when analyzing a model, check the Use risk preference function box and select either Constant risk aversion or Non-constant risk.

27.3.1 Constant risk aversion

If constant risk aversion is selected, you will be asked to supply a single value. Specifically, you will be shown a generic lottery in which you have a 0.5 probability of winning X and a 0.5 probability of losing one-half X , and asked to specify the largest value of X for which you would be willing to take part in the lottery. This value is used as the risk preference coefficient in the above formula.

The lottery might represent an investment in a biotech company which is about to get a judicial ruling on the validity of an important patent. If the ruling is favorable (0.5 probability), the investment will double in value; if unfavorable (0.5 probability), the investment will fall in value by 50%.

What is the most you would invest under these circumstances? This amount is referred to as your risk preference coefficient.

28. Using the Excel Module

The optional Excel Module enhances TreeAge Pro's ability to communicate with Excel. The major features of the Excel Module are:

1. Edit model inputs (variables, trackers, distributions and tables) in Excel.
2. Create a Tree Workbook to output the model inputs to Excel.
3. Output reports and graphs to Excel.
4. Access models programmatically through the Object Interface.

Features of the Excel Module

Each of these features are described within this chapter.



Dynamic bi-directional links with Excel are supported by the base TreeAge Pro product and do not require the Excel Module.

28.1 Edit model inputs in Excel

When you want to edit a large number of model inputs, it is often most efficient to do so via Excel. The process works as follows:

1. Export the desired inputs from TreeAge Pro to an Excel worksheet.
2. Edit the values in the Excel worksheet.
3. Import the new values back from Excel to the TreeAge Pro model.

Editing model inputs in Excel

This option is available for variables, trackers, distributions and tables. Each input is described in the next few sections. The Bilink tutorial example model TreeWorkbookTest.trex.





Updates from Excel are applied to the active model. Therefore, you must select the correct model in the Tree Diagram Editor before sending updates back from Excel to TreeAge Pro,

28.1.1 Edit variables in Excel

If you have the Excel Module licensed, the Variable Properties View toolbar's To Excel icon/function will be enabled.

To edit variables in Excel:

- Open the Variable Properties View.
- Click the Open in New Excel Spreadsheet toolbar icon.

Tree Properties Node Properties Variable Properties Variable Definitions				
Show Categories   type filter text <input type="text"/> clear				
Name	Description	Show in tree	Root Definition	Category
Variable01	Desc-Variable1	<input checked="" type="checkbox"/>	100	
Variable02	Desc-Variable2	<input checked="" type="checkbox"/>	200	
Variable03	Desc-Variable3	<input checked="" type="checkbox"/>	300	
Variable04	Desc-Variable4	<input checked="" type="checkbox"/>	400	
Variable05	Desc-Variable5	<input checked="" type="checkbox"/>	500	
Variable06	Desc-Variable6	<input checked="" type="checkbox"/>	600	
Variable07	Desc-Variable7	<input checked="" type="checkbox"/>	700	
Variable08	Desc-Variable8	<input checked="" type="checkbox"/>	800	
Variable09	Desc-Variable9	<input checked="" type="checkbox"/>	900	
Variable10	Desc-Variable10	<input checked="" type="checkbox"/>	1000	

Variable Properties View - To Excel function

The tree's variables are then exported to an Excel worksheet.



In the worksheet, the original variables are identified by hidden column I. This allows you to rename variables based on the current name. Hidden column H is used to determine which variables have been modified. We recommend that you do not modify the data in either hidden column.

Note that if you add new variables, you will not want to copy the original name in the hidden column. If you do, the existing variable will be replaced by the new variable.

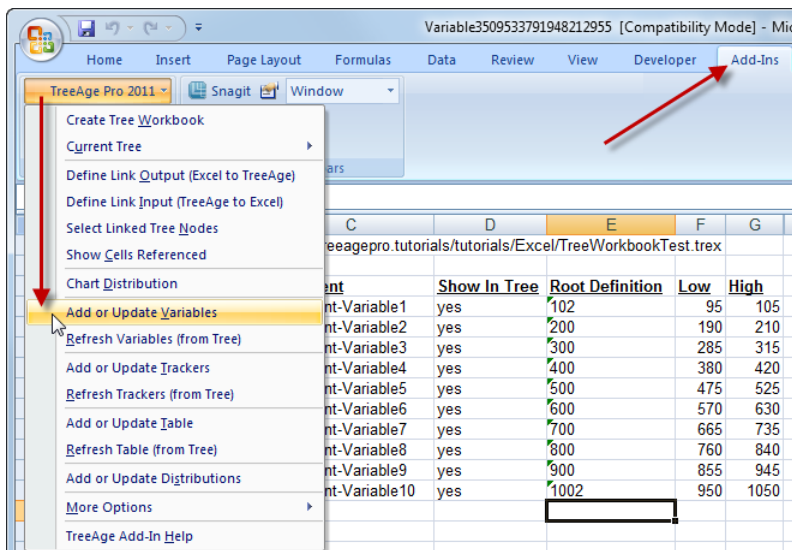
	A	B	C	D	E	F	G	I
1	Exported from: /resource/Examples/Tutorial%20Trees/Excel/TreeWorkbookTest.trex							
2								
3	Name	Description	Comment	Show In Tree	Root Definition	Low	High	Changed
4	Variable01	Desc-Variable1	Comment-Variable1	yes	100	95	105	0
5	Variable02	Desc-Variable2	Comment-Variable2	yes	200	190	210	0
6	Variable03	Desc-Variable3	Comment-Variable3	yes	300	285	315	0
7	Variable04	Desc-Variable4	Comment-Variable4	yes	400	380	420	0
8	Variable05	Desc-Variable5	Comment-Variable5	yes	500	475	525	0
9	Variable06	Desc-Variable6	Comment-Variable6	yes	600	570	630	0
10	Variable07	Desc-Variable7	Comment-Variable7	yes	700	665	735	0
11	Variable08	Desc-Variable8	Comment-Variable8	yes	800	760	840	0
12	Variable09	Desc-Variable9	Comment-Variable9	yes	900	855	945	0
13	Variable10	Desc-Variable10	Comment-Variable10	yes	1000	950	1050	0
14								

Variables worksheet in Excel

Within the worksheet, you can change the variable properties and the default definitions (at the root node).

To send the new properties/values back to TreeAge Pro:

- Select the correct tree in the Tree Diagram Editor.
- In Excel 2007 or later, choose the Add-ins ribbon then choose TreeAge Eclipse > Add or Update Variables from the Excel menu.
- In Excel 2003 or earlier, choose TreeAge Eclipse > Add or Update Variables from the Excel menu.



Update variables in Excel 2007

The variables in TreeAge Pro will be updated with the new properties/values from the Excel worksheet. There is no need to keep the Excel worksheet after the updates are complete as it can be regenerated at any time.

To refresh the data in the worksheet from the values in TreeAge Pro:

- In Excel 2007 or later, choose the Add-ins ribbon then choose TreeAge Eclipse > Refresh Variables from the Excel menu.
- In Excel 2003 or earlier, choose TreeAge Eclipse > Refresh Variables (from Tree) from the Excel menu.



Refresh will update the existing list of variables with the current values from TreeAge Pro. If you have added new variables within TreeAge Pro, please generate a new list via Edit in Excel from within TreeAge Pro.



If you export variables to Excel at a node other than the root node, you are asked whether to include an additional column of data for the variable definitions at the selected node. If you answer "yes", then the variable definitions at the selected node can also be updated and sent back to TreeAge Pro.

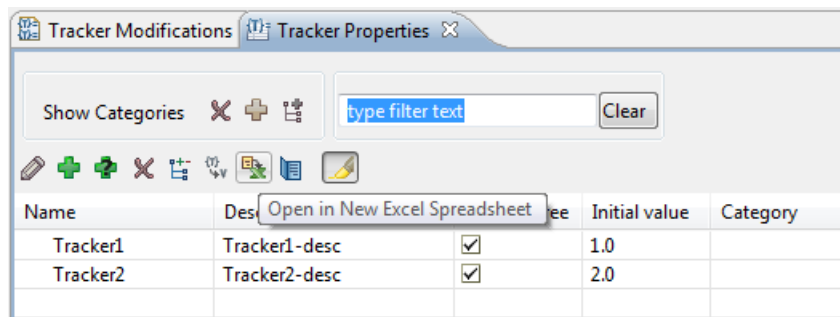
28.1.2 Edit trackers in Excel

Editing trackers in Excel works the same as editing variables, although tracker properties are different from variable properties.

If you have the Excel Module licensed, the Tracker Properties View toolbar's To Excel icon/function will be enabled.

To edit trackers in Excel:

- Open the Tracker Properties View.
- Click the Open in New Excel Spreadsheet toolbar icon (highlighted below).



Tracker Properties View - To Excel function

The tree's trackers are then exported to an Excel worksheet.

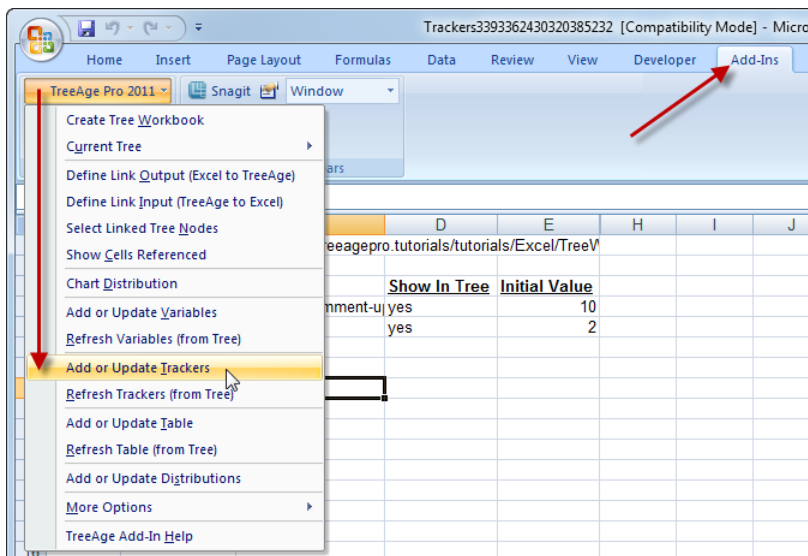
	A	B	C	D	E	G
1	Exported from: /resource/Examples/Tutorial%20Trees/Excel/TreeWorkbookTest					
2						
3	Name	Description	Comment	Show In Tree	Initial Value	Changed
4	Tracker1	Tracker1-desc	Tracker1-comment	yes	1	0
5	Tracker2	Tracker2-desc	Comment	yes	2	0
6						

Trackers worksheet in Excel

Within the worksheet, you can change the tracker properties and the initial values. Hidden columns are used in the same way as the variables worksheet. See note in prior section.

To send the new properties/values back to TreeAge Pro:

- Select the correct tree in the Tree Diagram Editor.
- In Excel 2007 or later, choose the Add-ins ribbon then choose TreeAge Eclipse > Add or Update Trackers from the Excel menu.
- In Excel 2003 or earlier, choose TreeAge Eclipse > Add or Update Trackers from the Excel menu.



Update trackers in Excel 2007

The trackers in TreeAge Pro will be updated with the new properties/values from the Excel worksheet. There is no need to keep the Excel worksheet after the updates are complete as it can be regenerated at any time.

To refresh the data in the worksheet from the values in TreeAge Pro:

- In Excel 2007 or later, choose the Add-ins ribbon then choose TreeAge Eclipse > Refresh Trackers from the Excel menu.
- In Excel 2003 or earlier, choose TreeAge Eclipse > Refresh Trackers from the Excel menu.



Refresh will update the existing list of variables with the current values from TreeAge Pro. If you have added new variables within TreeAge Pro, please generate a new list via Edit in Excel from within TreeAge Pro.



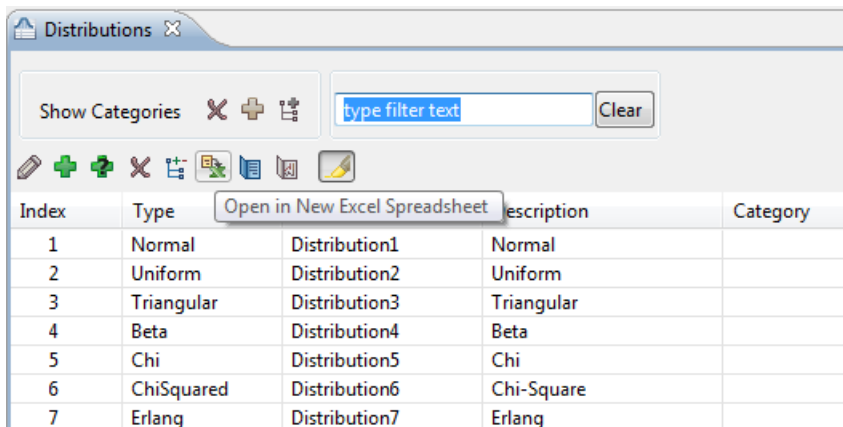
If you export trackers to Excel at a node other than the root node, you are asked whether to include an additional column of data for the tracker modifications at the selected node. If you answer "yes", then the tracker modifications at the selected node can also be updated and sent back to TreeAge Pro.

28.1.3 Edit distributions in Excel

If you have the Excel Module licensed, the Distributions View toolbar's To Excel icon/function will be enabled.

To edit distributions in Excel:

- Open the Distributions View.
- Click the Open in New Excel Spreadsheet toolbar icon (highlighted below).



Distributions View - To Excel function

The tree's distributions are then exported to an Excel worksheet. See note on hidden columns in Variables section above.

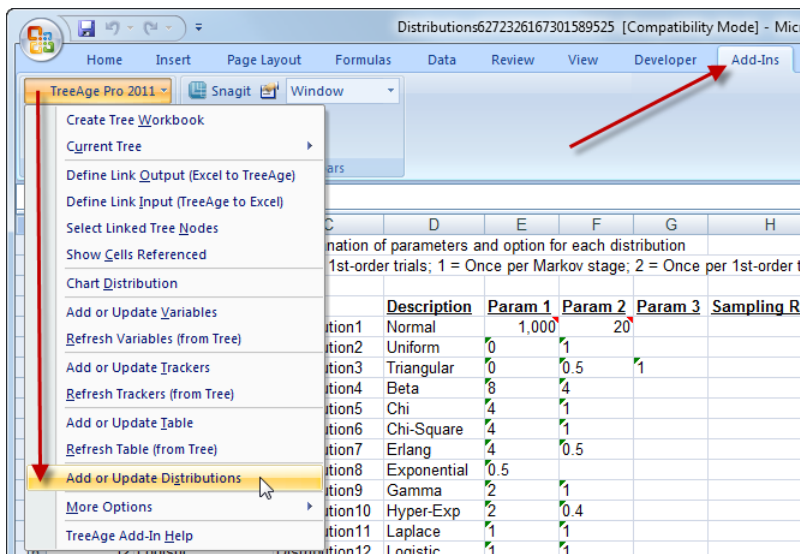
	A	B	C	D	E	F	G	H	L	M	N	O	P	Q
1	Exported from: /resource/Examples/Tutorial%20Trees/Excel/TreeWorkbookTest.tre													
2														
3	* see Help/Explanation column for explanation of parameters and option for each distribution													
4	* Sampling Rate: 0 = Once per group of 1st-order trials; 1 = Once per Markov stage; 2 = Once per 1st-order trial													
5														
6	Index	Type	Name	Description	Param 1	Param 2	Param 3	Sampling Rate	Changed	Help/Explanation				
7	1	Normal	Distribution1	Normal	1,000	75		0	0	0 Param 1 = mean; Param 2 = stddev;				
8	2	Uniform	Distribution2	Uniform	0	1		0	0	0 Param 1 = low; Param 2 = high;				
9	3	Triangular	Distribution3	Triangular	0	0.5	1	0	0	0 Param 1 = min; Param 2 = likeliest; Param 3 = max;				
10	4	Beta	Distribution4	Beta	8	4		0	0	0 Param 1 = n; Param 2 = r;				
11	5	Chi	Distribution5	Chi	4	1		0	0	0 Param 1 = n; Param 2 = sigma;				
12	6	ChiSquared	Distribution6	Chi-Square	4	1		0	0	0 Param 1 = n; Param 2 = sigma;				
13	7	Erlang	Distribution7	Erlang	4	0.5		0	0	0 Param 1 = k; Param 2 = lambda;				
14	8	Exponential	Distribution8	Exponential	0.5			0	0	0 Param 1 = lambda;				
15	9	Gamma	Distribution9	Gamma	2	1		0	0	0 Param 1 = alpha; Param 2 = lambda;				
16	10	HyperExponential	Distribution10	Hyper-Exp	2	0.4		0	0	0 Param 1 = lambda; Param 2 = p;				
17	11	Laplace	Distribution11	Laplace	1	1		0	0	0 Param 1 = a; Param 2 = b;				
18	12	Logistic	Distribution12	Logistic	1	1		0	0	0 Param 1 = a; Param 2 = b;				
19	13	LogNormal	Distribution13	Log-Normal	5	1		0	0	0 Param 1 = umeanoflogs; Param 2 = sigmastddevoflogs;				
20	14	Maxwell	Distribution14	Maxwell	5			0	0	0 Param 1 = alpha;				
21	15	Rayleigh	Distribution15	Rayleigh	1			0	0	0 Param 1 = alpha;				
22	16	Weibull	Distribution16	Weibull	1	2		0	0	0 Param 1 = scalefactor; Param 2 = shape;				
23	17	Poisson	Distribution17	Poisson	10			0	0	0 Param 1 = lambda;				
24	18	Binomial	Distribution18	Binomial	0.1	20		0	0	0 Param 1 = probability; Param 2 = trials;				
25	19	Fractile	Distribution19	Bin-50-90	1	5	9	0	0	0 Param 1 = 10; Param 2 = 50; Param 3 = 90;				

Distributions worksheet in Excel

Within the worksheet, you can change the distribution properties and parameters. Note that the Help/Explanation column provides details on how the parameters are used for the appropriate distribution type.

To send the new properties/parameters back to TreeAge Pro:

- Select the correct tree in the Tree Diagram Editor.
- In Excel 2007 or later, choose the Add-ins ribbon then choose TreeAge Eclipse > Add or Update Distributions from the Excel menu.
- In Excel 2003 or earlier, choose TreeAge Eclipse > Add or Update Distributions from the Excel menu.



Update trackers in Excel 2007

The distributions in TreeAge Pro will be updated with the new properties/parameters from the Excel worksheet. There is no need to keep the Excel worksheet after the updates are complete as it can be regenerated at any time.



A few distribution types have additional options that are not updated through the Excel module (e.g., Fractile Gaussian vs. Swanson). These options must be edited in the Add/Change Distribution dialog.

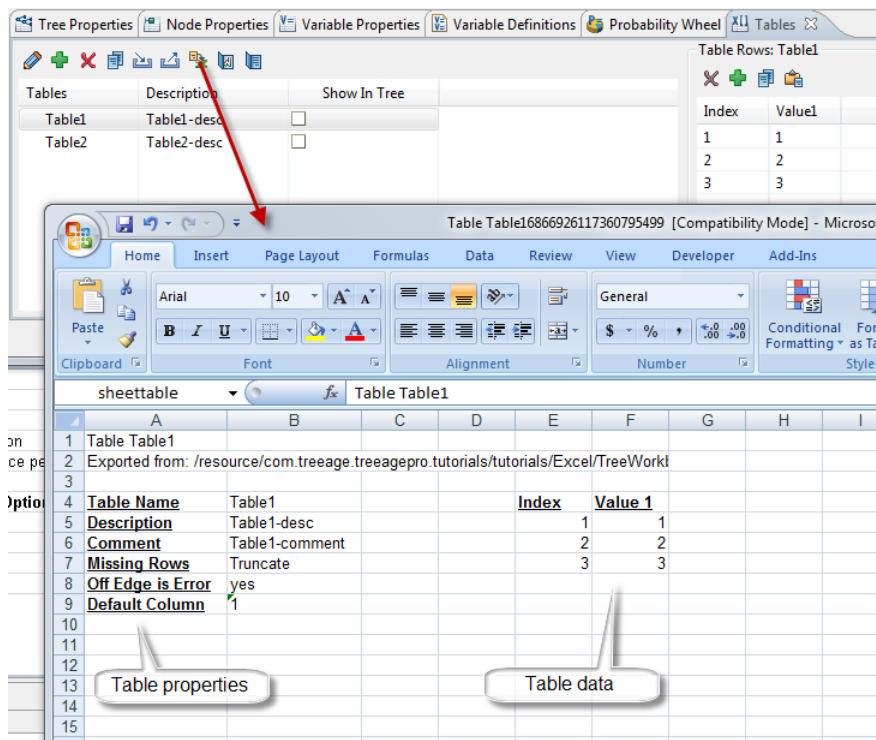
28.1.4 Edit tables in Excel

Variables, trackers and distributions are edited within a list. However, tables are edited individually in Excel since they contain both properties and data.

If you have the Excel Module licensed, the Tables View toolbar's Open in New Excel Spreadsheet icon/function will be enabled.

To edit a table in Excel:

- Open the Tables View.
- Select a single table.
- Click the To Excel toolbar icon (highlighted below), which exports the table to an Excel worksheet.



Tables View - To Excel Function

Within the worksheet, you can change the table properties and data. Be careful not to change the structure and/or location of the data in the worksheet.

To send the new properties/data back to TreeAge Pro:

- Select the correct tree in the Tree Diagram Editor.
- In Excel 2007 or later, choose the Add-ins ribbon then choose TreeAge Eclipse > Add or Update Distributions from the Excel menu.
- In Excel 2003 or earlier, choose TreeAge Eclipse > Add or Update Distributions from the Excel menu.

To refresh the data in the worksheet from the values in TreeAge Pro:

- In Excel 2007 or later, choose the Add-ins ribbon then choose TreeAge Eclipse > Refresh Trackers from the Excel menu.
- In Excel 2003 or earlier, choose TreeAge Eclipse > Refresh Trackers from the Excel menu.



Note that you can add columns to the table in Excel by adding data to the right of the existing columns. Value columns can have custom headings, but the Index column heading should not be changed.

28.2 Tree Workbook

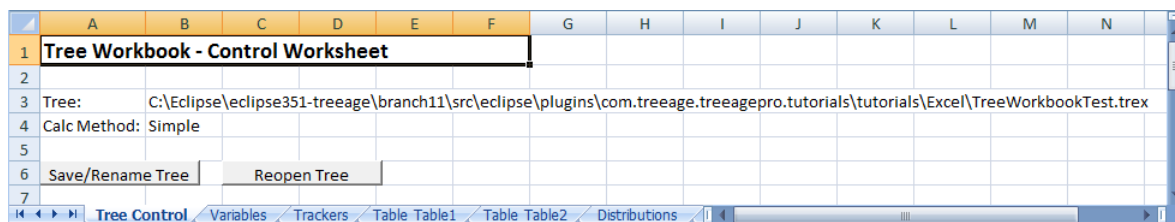
You can quickly export the model inputs described in the previous sections (variables, trackers, distributions and tables) via a Tree Workbook.

To create a Tree Workbook:

- In Excel 2007 or later, choose the Add-ins ribbon then choose TreeAge Eclipse > Create Tree Workbook from the Excel menu.
- In Excel 2003 or earlier, choose TreeAge > Create Tree Workbook from the Excel menu.

A new Excel workbook will be created with a separate worksheet for each model input (variable list, tracker list, distribution list and each table). These model input worksheets are the same as described previously for variables, trackers, distributions and tables. Additional worksheets are described within this section.

The Tree Control worksheet is used to either open or save the model associated with the Tree Workbook. You can change the name of the tree and save the model in a separate document.



Tree Workbook



If you have different data scenarios you want to save for the same model. You can save multiple Tree Workbooks for the model. Then, you can use the Tree Workbook to update the model's data for that specific data scenario.

28.3 TreeAge Pro Object Interface

This interface is described in the TreeAge Pro Object Interface Chapter.

28.4 Exporting analysis output

The Excel module also allows exporting analysis outputs to Excel. The Open in New Excel Spreadsheet icon is frequently enabled for graphical and text output.



Open in New Excel Spreadsheet icon

When this icon is enabled at the top right of the output, you can use it to export the results to Excel.

29. Using the TreeAge Pro Object Interface

The Excel Module (included with TreeAge Pro Excel and TreeAge Pro Suite) provides a powerful scripting interface that can be used to open, update and analyze models via program code.



The Object Interface for TreeAge Pro is different from previous versions of TreeAge Pro. You can continue to use the older version of TreeAge Pro as you choose to modify the code which accesses TreeAge Pro.

29.1 Java vs. Other Programming Languages

You can open, update and analyze models via program code written in virtually any language, including:

- Java
- Visual Basic in an Excel macro
- Visual Basic in other environments
- C++
- etc.

TreeAge Pro is written in Java, so the Java programming language provides the most transparent look at the Object Interface via the Java package `com.treeage.treeagepro.oi`. This package includes the model element and analysis objects, which makes coding in Java easier because the Java IDE can "see" the TreeAge Pro objects and provides guidance on programming syntax.

However, many TreeAge Pro customers use the Object Interface through other programs, frequently via Visual Basic in an Excel macro. Languages other than Java connect to the Object Interface through the ActiveX Java package `com.treeage.treeagepro.oi.activex`. The ActiveX Java package allows you to instantiate the `TreeAgeProApplication` object. The `TreeAgeProApplication` object then provides access to instantiate other objects (trees, nodes, etc. from `com.treeage.treeagepro.oi`).

The API documentation describes the syntax for both Java packages.

29.2 Object Interface API Documentation

The Object Interface's Application Programming Interface (API) is described via standard javadoc documentation, which provides documentation for all objects and methods supported by the Object Interface.

A few of the primary application and modeling objects are highlighted in the following table. These and other objects are documented in the API documentation.

Object	Description
<code>TreeAgeProApplication</code>	A reference to the TreeAge Pro application. This is the first object you must create to start using the Object Interface when not using Java.

Object	Description
Tree	Standard tree model created in TreeAge Pro.
Node	A specific node within the tree.
Variable	A variable within a tree. Use this object to read/update variable properties.
VariableDefinition	A variable definition within a tree at a specific node.
Tracker	A tracker within a tree. Use this object to read/update tracker properties.
TrackerModification	A tracker modification within a tree at a specific node.
Table	A table within a tree. Use this object to read/update table properties.
TableRow	A row of data within a table.
Distribution	A distribution within a tree. Use this object to read/update the distribution's type, parameters and properties.

Primary modeling objects

The Object Interface Additional also includes objects that support analysis. Most analyses are run on the Tree object. However, several types of analyses require objects for handling inputs/parameters and outputs.

A few of the primary application and modeling objects are highlighted in the following table. These and other objects are documented in the API documentation.

Object	Description
Report	Output from any analysis. The type of analysis executed determines the format of the data within the object. Some Report objects can be used to create other Report objects to generate secondary analysis outputs (i.e., graphs from Monte Carlo simulation output). Analysis parameters are passed via a hashmap of named values prior to generating the report.
Graph	Graphical output from a Report object.
TextReport	Tabular text and numeric output from a Report object.

Analysis objects

29.3 Connecting to the Object Interface via Java

When using Java code to connect to the Object Interface, you must create the appropriate file structure to be able to "build" your Java project. This requires some experience with creating a Java build environment.

To avoid the need to create an appropriate development environment, you can simply copy the full structure of the Object Interface Example project (described below) to another location on your computer and network. With the same file structure, you will be able to create Java source code in the folder `../src/com/treeage/treeagepro/tutorials/`, placing your new source files alongside the example file `UseObjectInterface.java`.

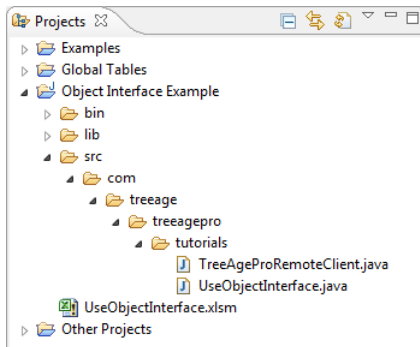


Make sure you do not make changes directly in the Object Interface Example project as this project could be overwritten when new software is released.

If a new version of the Object Interface Example project is released, copy the appropriate files to your project. Be sure not to overwrite any of your own source code.

29.3.1 Sample Java code

Sample Java code is provided in the Object Interface Example project installed with your software. You can access the appropriate file through the Projects View (see below).



Object Interface Example project

The `UseObjectInterface.java` source file contains sample Java code that opens, edits and analyzes sample trees. You can use this file as a starting point for creating your own Java source code.



If you choose to create your own source code using this example, be sure to save the file using a different filename. In fact, we recommend copying the entire project to a separate location. Otherwise, your source could be overwritten if an updated version of the example code is released.

This section will introduce you to the sample Java code. However, the Java code itself contains comments that serve as the primary documentation.

The `UseObjectInterface.java` file contains a Java class of the same name. That class runs as a Java application based on the existence of the method *main*. The *main* method instantiates the class, which immediately calls the class constructor *UseObjectInterface*, which contains the main processing steps. Additional methods provide the details on opening, modifying and analyzing trees.

```

/**
 * Constructor for class UseObjectInterface.
 *
 * Contains main tree processing.
 */
private UseObjectInterface() {

    com.treeage.treeagepro.oi.Tree sampleTree;
    com.treeage.treeagepro.oi.Tree ceSimTree;

    // Connects locally to TP2011. For remote use see TreeAgeProApplication
    // constructors
    // such as TreeAgeProApplication(host).
    app = new TreeAgeProApplication();
    if (!app.isValid()) {
        System.out.println("Cannot find TP2011 application running locally.");
        return;
    }

    try {
        System.out.println("Workspace path: " + app.getWorkspacePath());
        // Open a sample tree. The path can be either absolute or relative
        // to workspace root folder.
        sampleTree = this.openTree(sampleTreeOrig);

        // Save under a new name. The path can be either absolute or
        // relative to workspace root folder.
        sampleTree.saveAs(sampleTreeNew);

        // Output tree characteristics
        outputTreeInfo(sampleTree);

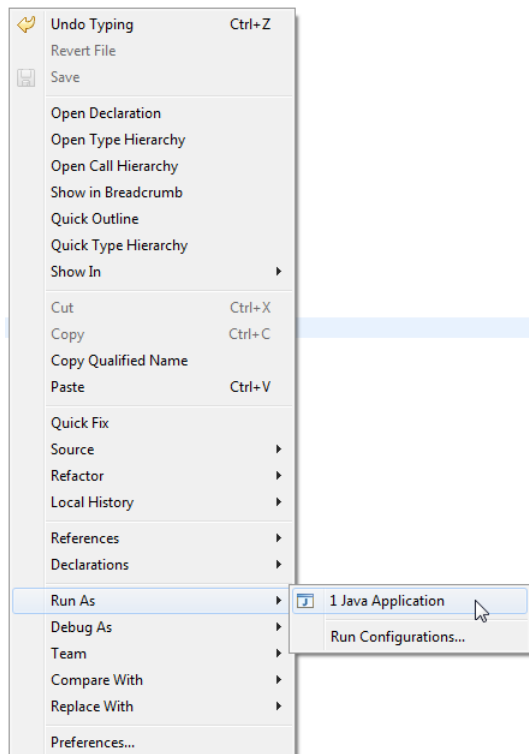
        // Read/update variables.
        updateVariable(sampleTree);

        // Analyze a simple tree and report results.
        analyzeTree(sampleTree);
    }
}

```

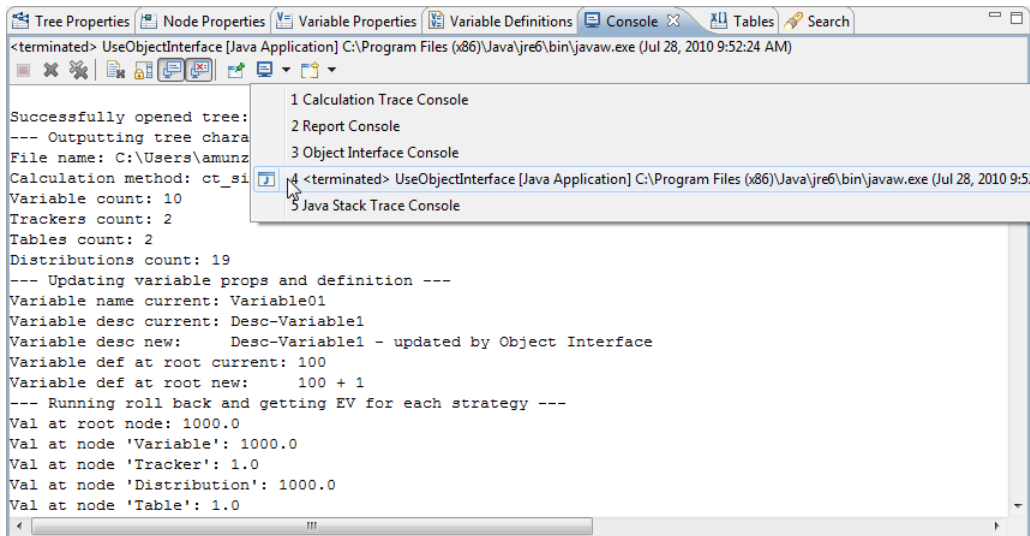
Sample Java code

To execute the code, right-click in the Java code document and select Run As > Java Application from the menu.



Execute Java code

As the sample code executes, it writes a status output to the Java output Console. See below.

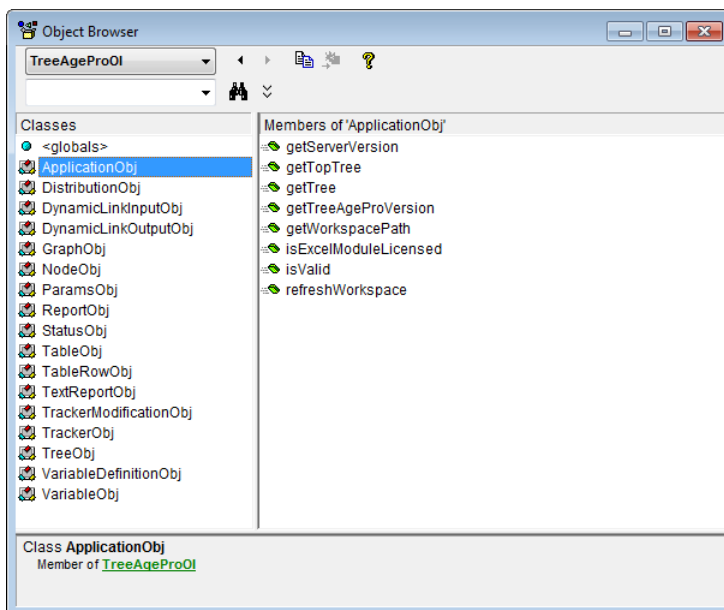


Java Console output

29.4 Connecting to the Object Interface via ActiveX

Scripting languages communicate with TreeAge Pro via Java Remote Method Invocation (RMI). When TreeAge Pro starts up it starts a service on port 1099 (or slightly higher) which listens for ActiveX requests.

The non-Java objects are instantiated via the library TreeAgeProOI. This library must be referenced for your code to access the TreeAge Pro Object Interface objects. Below is a picture of the library objects as seen from the Excel Visual Basic Editor's Object Browser.



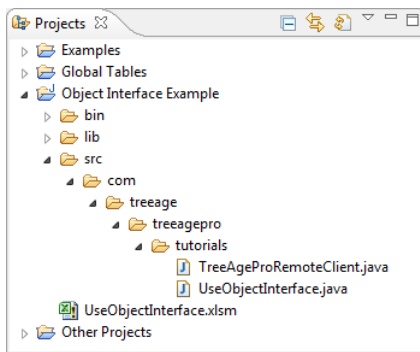
TreeAgeProOI Objects in Excel VB Object Browser

You can use these library objects to open, modify and analyze trees within Excel or other programming environments. Even when not working in Java, refer to the Java API documentation for object/method syntax.

Since ActiveX is registered on the computer, there are no requirements for a build environment as there are for Java projects. You can save the code anywhere on your computer or network.

29.4.1 Sample Visual Basic code

Sample Visual Basic (VB) code is provided in the Excel document "Use Object Interface.xls" in the TreeAge Pro Interface tutorial examples project. See below.



VB code sample Excel document

The UseObjectInterface.xlsm document contains sample VB code that opens, edits and analyzes sample trees. You can use this file as a starting point for creating your own VB source code.



If you choose to create your own source code using this example, be sure to save the file using a different filename in a different location. Otherwise, your source could be overwritten if an updated version of the example code is released.

This section will introduce you to the sample VB code. However, the VB code itself contains comments that serve as the primary documentation.

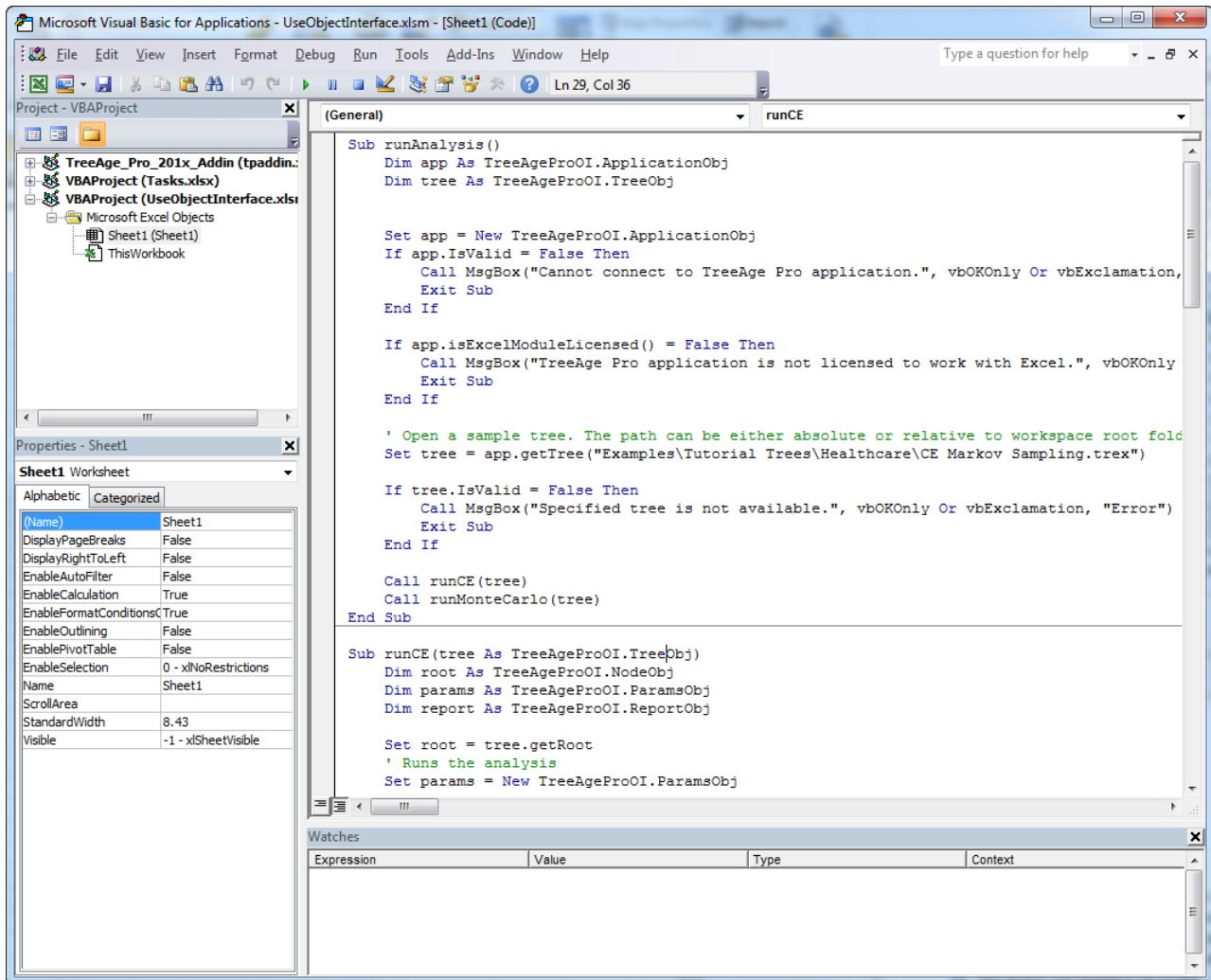
To see the VB code in Excel 2007 or later:

- Click on the Office button and then click the Excel Options button.
- In the Popular category, make sure the Show Developer tab in the ribbon box is checked.
- Select the Developer tab in the ribbon.
- Click the Visual Basic button.

To see the VB code in Excel 2003 or earlier:

- Choose Tools > Macros > Visual Basic Editor from the menu.

The VB code has a primary macro called TestObjectInterface, which contains the main processing steps. Additional methods provide the details on opening, modifying and analyzing trees.



Sample code and output in Excel's VB editor

30. The TreeAge Pro Player

TreeAge Pro 2011 R2.0 introduced a new feature called the TreeAge Pro Player. The Player allows you to share a model with an individual who does not have a license for TreeAge Pro. A consultant might use this feature to share the model to his/her client.

In order to use the Player, the modeler must first create a Player Interface for the model, resulting in a password protected Player Model with the *.trvx extension. The modeler determines which input variables can be changed and which analyses can be executed. The modeler also determines whether or not the player should be able to display the model structure.

The Player Model can be opened with TreeAge Pro, but functions are limited to those exposed by the model interface. Supplying the password will allow you to use an expanded set of functions.

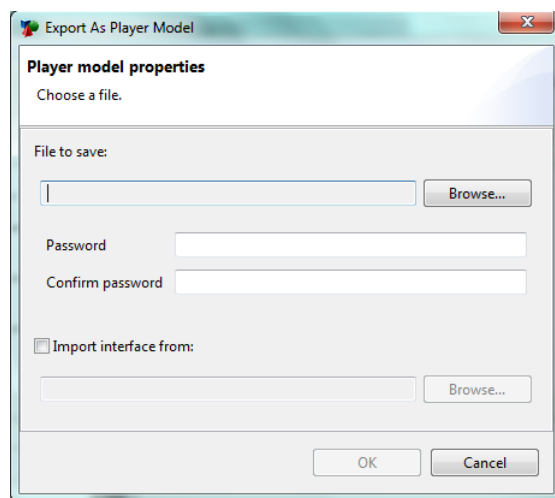
30.1 Creating a Player Model

You create a Player Model by exporting a model in the player format.

To create a Player Model:

1. Save the latest changes to the model.
2. Choose File > Import/Export > Export as Player Model from the menu.
3. Choose the file name and location in the dialog box.
4. Enter the password associated with the Player Model twice.

The model will be exported as a Player Model will be saved with the *.trvx extension.



Export as Player Model Dialog



You cannot change the structure or values within a model within a Player Model. You must return to the original model file to make such changes. However, you can then create a new Player Model and import the Model Interface elements from an existing Player Model.

If you are modifying a model for which an interface had already been created, you have the option of importing the interface from another Player Model. To do so, follow the instructions above plus the following steps:

1. Check the Import interface box.
2. Select an existing Player Model (*.trvx) file.

The resulting Player Model will include the model structure and details from the model that was exported as well as the interface details from the prior Player Model.



If any variable names or stored analysis names from the imported interface are not found in the exported model, they will not be usable in the new interface. They will be displayed in red and can be deleted but cannot be used. This can happen if a variable or stored analysis is renamed since a prior interface was created.

30.2 Creating the Player Model Interface

Once the Player Model has been exported, you then need to create a Model Interface to provide access to the parameters and analyses. When you open the Player Model, TreeAge Pro will switch to the Player perspective. The Player Perspective limits access to nearly all of the application views used for editing the model.

The Player Perspective includes the following Views:

- Player Model - for using an existing Model Interface.
- Player Design - for creating a Model Interface.
- Tree Diagram Editor - for viewing the model (if allowed by the interface)
- Console - for viewing system output from analysis runs
- Error Log - for viewing system errors
- Projects - for accessing model files

If no interface has been created, you will only able to view the model.

To create a Player Model Interface:

1. Open the Player Model.
2. Choose Edit > Enter the Designer Password.
3. Enter the password used when creating the Player Model.



If you do not have an authorized version of TreeAge Pro, you will not be able to enter the Designer Password.

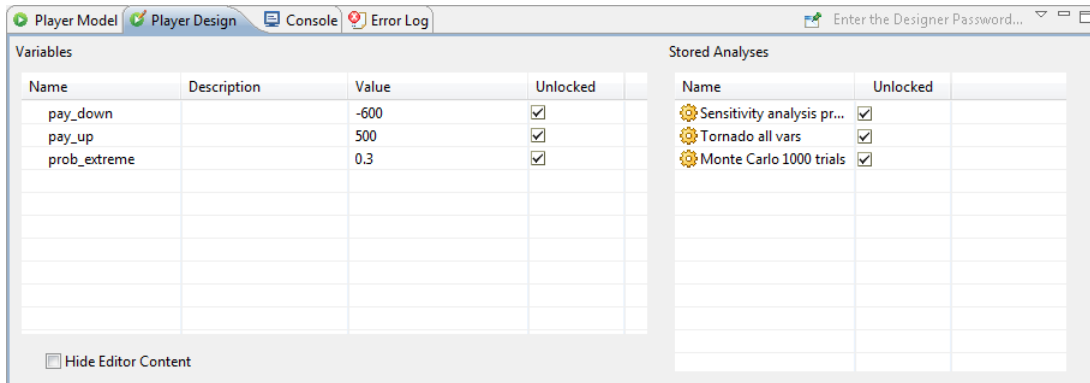
The Player Design View will become enabled. You create the interface within this view.

There are several components of the interface:

- Variable parameters that can be updated.
- Stored analyses that can be executed.
- A setting to determine whether the interface should show the model structure or not.

Interface Components

The figure below shows the Player Design View with all of the variables exposed, all stored analyses exposed and the model structure exposed to the interface. This interface was created from the Special Features tutorial example model Three Vars with Stored Analyses.



Player Design View

Variables

A model will have many variables that serve several functions in the model. Some variables are model parameters with a numeric definition at the root node. Only variables such as these should be enabled in the Model Interface, and probably only a subset of those variables.

Check the Unlocked box next to each variable you wish to expose in the Model Interface. It will be helpful to the person using the Model Interface if the variable has a clear name and description. You can also enter additional information in the Comments field.

Stored Analyses

Within the Model Interface, the regular Analysis menu will not be available. Instead, the modeler must create a stored analysis for each analysis they wish to expose in the Model Interface. Comments can be added to the stored analysis to help guide the user of the Model Interface.

Check the Unlocked box next to every stored analysis you wish to expose in the Model Interface.



You can restrict the secondary output options available within the standard Monte Carlo simulation output to make it easier for the Player Model recipient to find the output of interest.

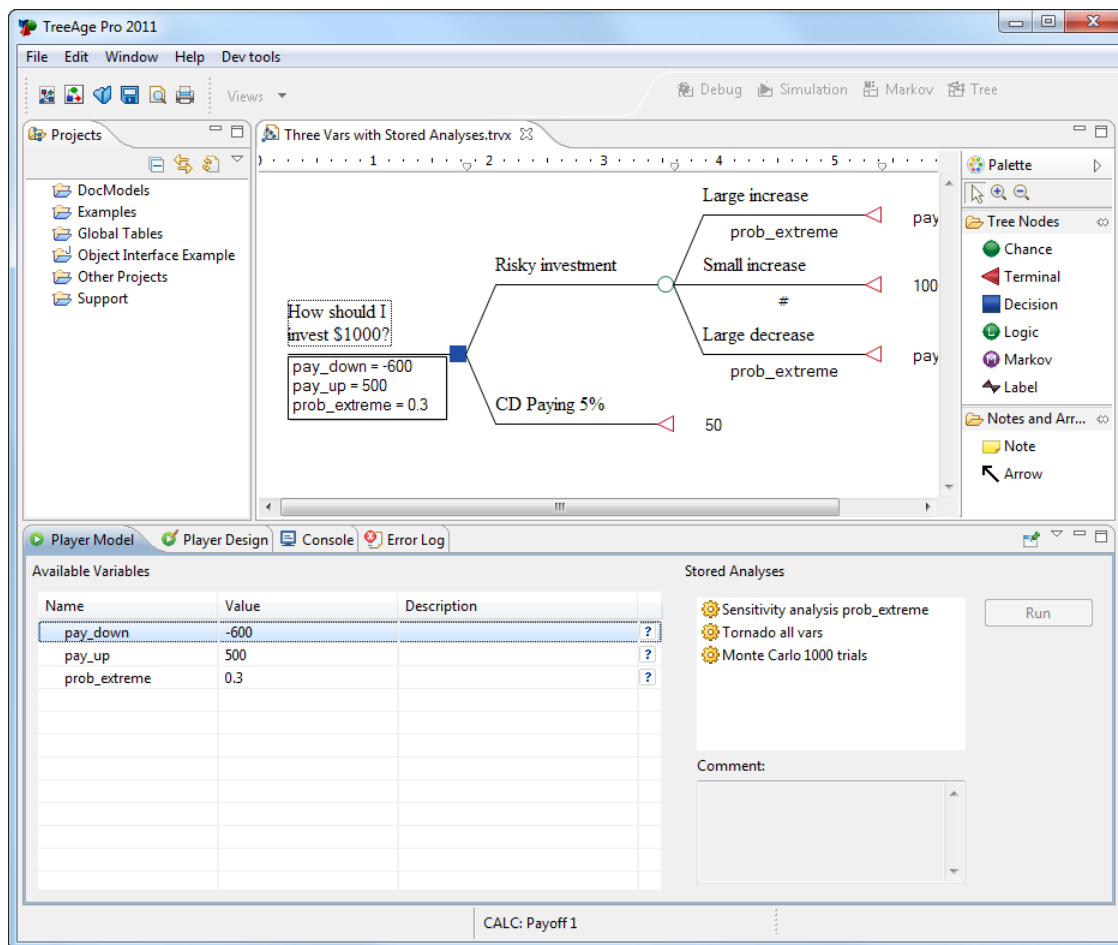
Exposing the Model Structure

The modeler may or may not want the Model Interface to expose the structure of the model. This is controlled by checking or unchecking the box in the Player Design View.

30.3 Using the Model Interface

Once the Model Interface is complete, the Player Model can be opened and analyzed using TreeAge Pro even if the software is not authorized. When the Player Model is opened, the Model Interface will present the specific variable inputs that can be modified, allowing the user to change certain model assumptions. Variable descriptions and comments in the original model can be used to guide someone using the interface, including specifying valid values.

The Model Interface will also show the specific analyses that can be run. The user can run any of the specified analyses and the appropriate output will be presented.



Player View

31. Building and Analyzing Cost-Effectiveness Models

The next three chapters provide information on modifying existing decision trees for cost-effectiveness analysis. Cost-effectiveness analysis, sensitivity analysis, and Monte Carlo simulation are also covered in detail. The three chapters related to these topics are:

1. Building and Analyzing Cost-Effectiveness Models (this chapter)
2. Cost-Effectiveness Modeling and Analysis Options
3. Cost-Effectiveness Simulation Reports and Graphs

Cost-effectiveness chapters

Please note that you must have purchased the Healthcare Module to take advantage of the cost-effectiveness features.

Healthcare decisions often take into account differences in both cost and effectiveness between competing treatments, technologies, or strategies. This chapter covers preparing an existing TreeAge Pro decision tree for cost-effectiveness calculations, and performing and interpreting baseline cost-effectiveness analysis.

The Cost-Effectiveness Modeling and Analysis Options provide information on intermediate and advanced cost-effectiveness modeling and analysis options.

31.1 Before you begin

Cost-effectiveness analysis (CEA) is a collection of methods used by health economists and policy researchers to evaluate policy recommendations on the basis of two different attribute scales (i.e., cost and quality-adjusted life expectancy).

The Healthcare Module (included with TreeAge Pro Healthcare and TreeAge Pro Suite) facilitates cost-effectiveness analysis and survival/Markov modeling and simulation — both so-called *early* models which rely heavily on rough guesses about outcomes and parameters, as well as *evidence-based* models that summarize extensive research and meta-analysis.

The basis of CEA in a decision tree is the calculation of expected values for each strategy at a decision node. TreeAge Pro then creates a CEA table, in which the strategies are listed in order of increasing cost, and calculates incremental cost and effectiveness values for the neighboring pairs of options. This is used to determine conditions of *dominance* and to calculate *incremental cost-effectiveness ratios* (ICERs), as described in this chapter.

For more background, users are strongly encouraged to consult some of the many references on medical decision making, for instance:

- Periodicals including *Medical Decision Making* (Sage Science Press) and *Value in Health* (Blackwell Publishing).

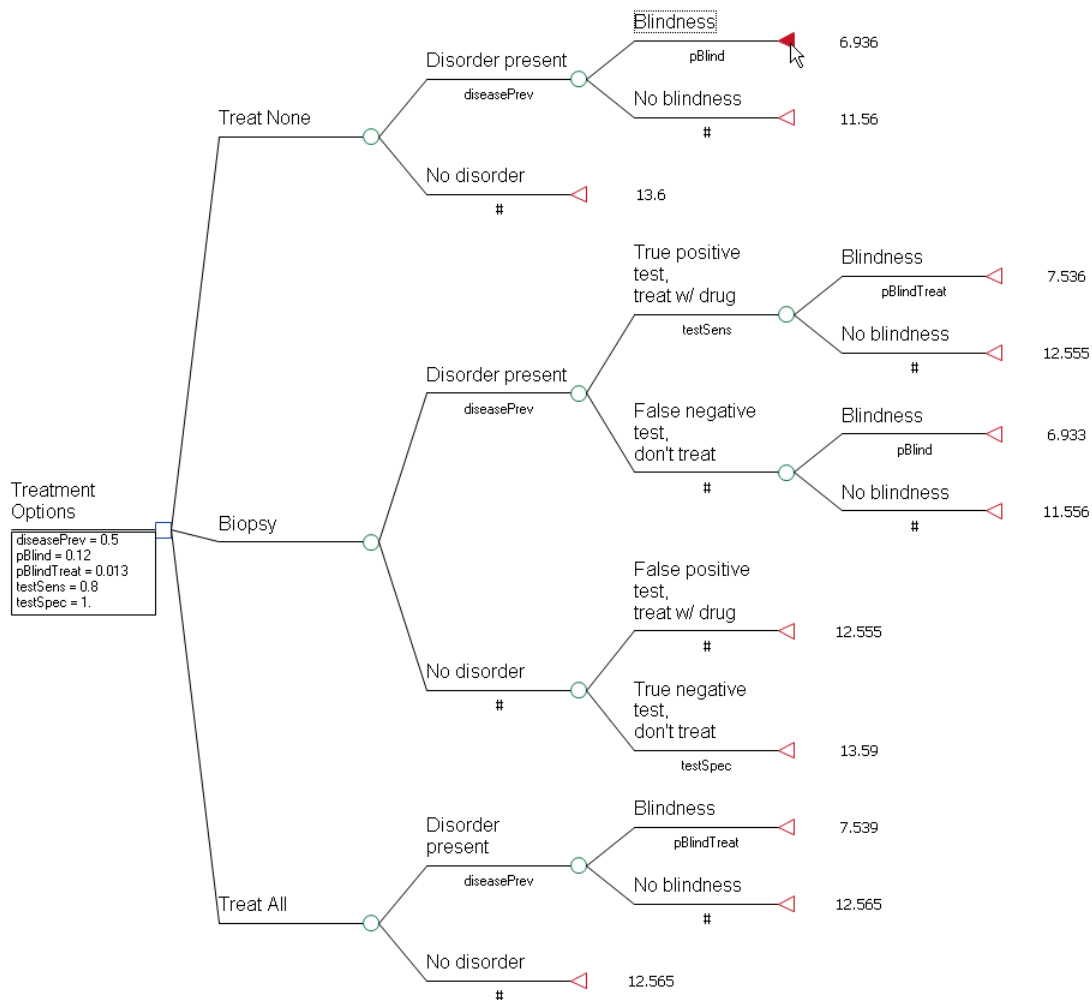
- *Cost-effectiveness in Health and Medicine*, Gold, Siegel, Russell, and Weinstein, eds. (1996), New York: Oxford Univ. Press.
- *Decision Making in Health and Medicine*, Hunink, and Glasziou (2001), Cambridge University.
- *Decision Modeling for Health Economic Evaluation*, Briggs, Claxton, and Sculpher (2007), Oxford Univ. Press.
- *Designing and Conducting Cost-Effectiveness Analyses in Medicine and Health Care, 2nd Ed.*, Muennig. (2007), New York: Oxford Univ. Press.
- *Medical Decision Making*, Sox, et al. (1988), Boston: Butterworth-Heinemann.
- *Meta-Analysis, Decision Analysis, and Cost-Effectiveness Analysis*, Petitti, (1994), New York: Oxford Univ. Press.
- *Methods for the Economic Evaluation of Health Care Programmes, 3rd Ed.*, Drummond, et al (2005), New York: Oxford Univ. Press.

Cost-effectiveness analysis references

31.2 Preparing a tree for cost-effectiveness calculations

This chapter uses an example decision tree that already has costs and effectiveness values in two payoffs. The tree is an adaptation of a model described in other decision analysis tutorials.

Open the Health Care tutorial example model "Blindness Prevention - before CE changes".



Blindness prevention model

The model deals with a hypothetical population presenting clinical signs of a possible, but not certain, early-stage autoimmune disorder. If the condition is present, and if it progresses, blindness will result. An imperfect test (biopsy, with the possibility of false negatives) can help determine whether an individual has the disorder.

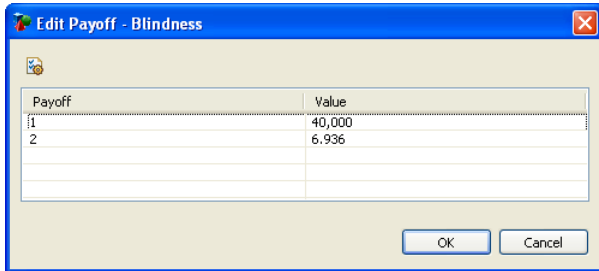
The model presumes that a reasonably effective and inexpensive therapy exists which lowers the probability of progression to blindness. To keep the example tree small, the possibility of side effects of the therapy is not modeled with a chance node like the other uncertainties. Instead, the side effects are already factored into the costs and life expectancy at the end of each path including treatment.

31.2.1 The Cost-Effectiveness calculation method

Before making any changes to the Blindness Prevention tree, take a moment to examine the assignment of payoffs at a terminal node.

To examine a payoff:

- Right-click on the Blindness terminal node in the Treat None subtree (marked by the pointer in the figure above) and select Edit Payoffs from the context menu. This will open the Edit Payoff Dialog for that node.



Edit payoff dialog

The numeric effectiveness payoffs displayed on the face of the tree, measuring quality-adjusted life expectancy (QALYs), are entered in payoff 2. In addition, every terminal node also has a numeric cost payoff assigned in payoff 1. The cost payoffs are not displayed, however, because the tree's calculation preferences are set to Simple calculations, using only payoff #2.



Note on regional (European) numeric settings:

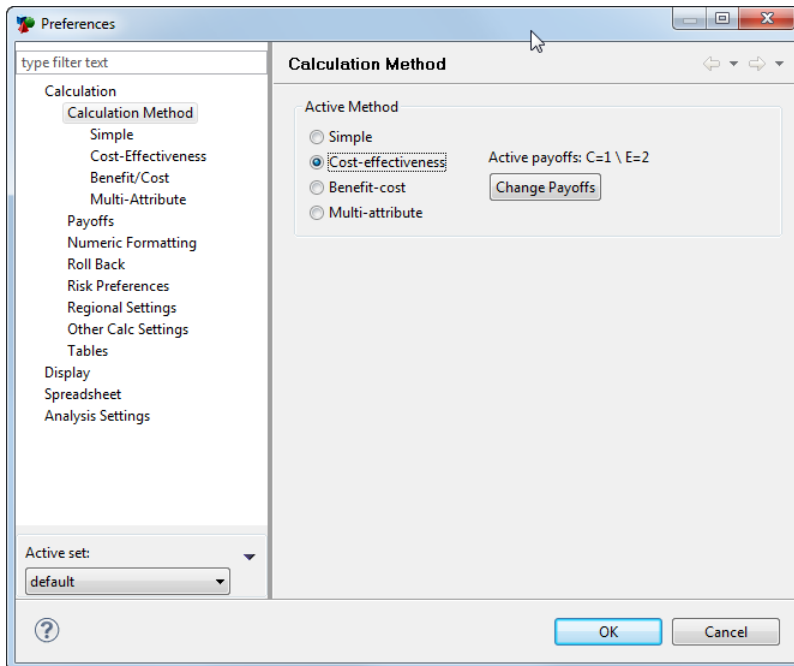
If your computer is set up to use commas (",") to represent decimals, rather than periods ((".")), you should enter numbers in TreeAge Pro in this fashion, just as you would in a spreadsheet or calculator (even in example trees where numbers already appear using period decimals). However, if you need to setup a tree for use under different regional numeric settings, TreeAge Pro also has preference settings that enable a tree to override a computer's regional settings or reverse the usage of decimals. Refer to the Tree Calculation Methods and Preferences Chapter for details.

Before you can perform cost-effectiveness analysis, the tree's calculation method preferences must be set up correctly.

You can do this using the Configure Model wizard or you can do this manually. In this section, we describe how to set Tree Preferences manually, but the same steps can be carried out using the Configure Model wizard.

To prepare a tree for cost-effectiveness calculations:

- Close the Edit Payoff Dialog.
- Choose Tree > Tree Preferences from the menu or press the *F11* key to open the Tree Preferences Dialog.
- Navigate to the category Calculation > Calculation Method.
- Select the option Cost-Effectiveness.



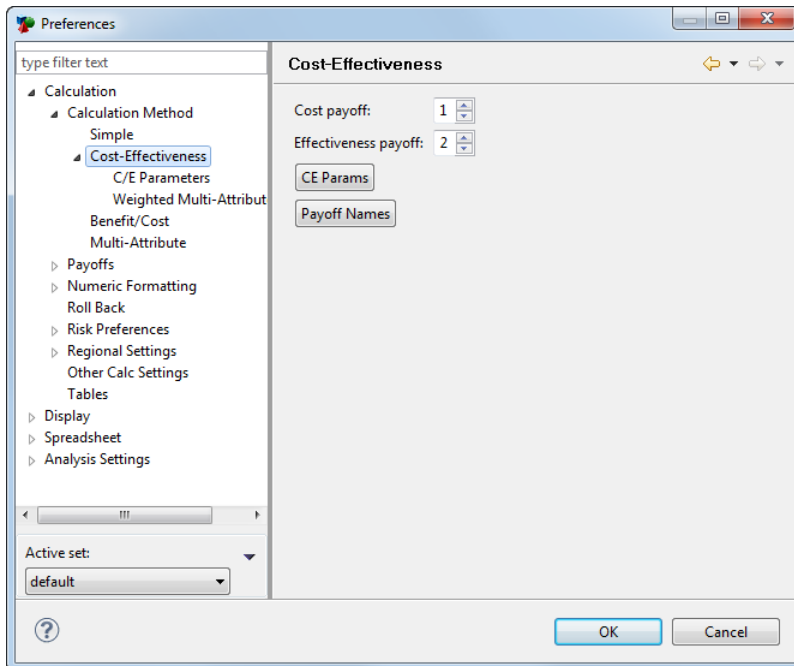
Calculation Method - Cost-Effectiveness

This will enable the special analyses and reports described in this and following chapters. Note that the tree can be changed back to Simple calculations at any time.

To review additional preferences related to cost-effectiveness analysis:

- In the Tree Preferences Dialog, navigate to the category Calculation > Calculation Method > Cost-Effectiveness.
- In this category, you can change the active payoffs for cost and effectiveness (not necessary for this tree).

By default, the Cost-Effectiveness calculation method uses payoff 1 for costs, and payoff 2 for effectiveness; this can be changed. However, in the Blindness Prevention tree, costs are already in payoff 1 and effectiveness values are in payoff 2.



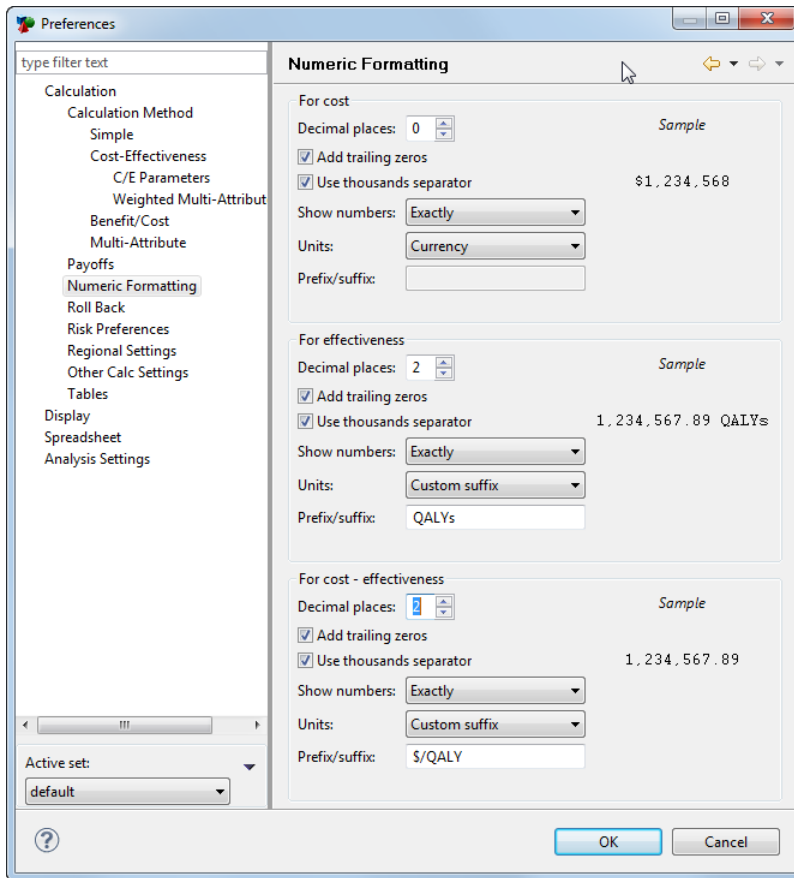
Cost-Effectiveness Payoffs

To set the numeric formatting for calculated costs, effectiveness values, and CE ratios:

- In the Tree Preferences Dialog, navigate to the category Calculation > Numeric Formatting.
- Click OK to save all changes to the Tree Preferences.

The tree stores separate numeric formatting preferences for cost and effectiveness (as well as the other 7 payoff sets), and another group of settings for ratios.

The graphic below illustrates some possible settings for these three sets of numeric formatting used in cost-effectiveness analysis.



Cost-Effectiveness Numeric Formatting

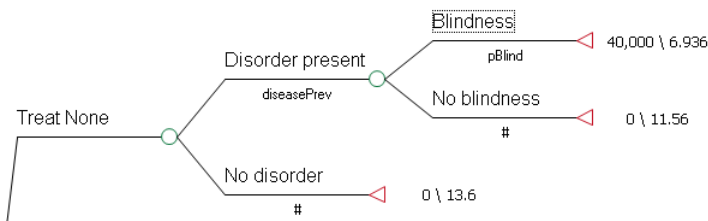


The Blindness Prevention model saved with the Tree Preference changes described in this section is in the Health Care tutorial example model "Blindness Prevention - after CE changes.trex". You can open the updated model directly to run the analyses described below if you want to skip the Tree Preferences changes.

If you plan to create more cost-effectiveness trees, you can use this tree's calculation method, numeric formatting, and other preferences as the defaults when creating new trees.

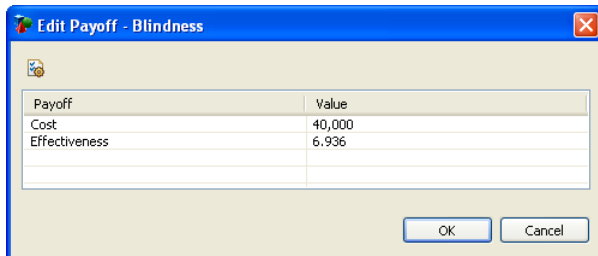
The next chapter covers additional special cost-effectiveness preferences, for example: setting a threshold ICER for roll back; inverting effectiveness calculations (if lower values are preferred); or specifying costs using a weighted combination of multiple payoffs.

With the CE calculation method active, the tree displays both the cost and effectiveness payoff expressions for each visible terminal node. A forward slash ("/") sign is used to visually separate the two payoffs; however, it *does not* mean that cost will be divided by effectiveness during roll back or other tree calculations.



Portion of Blindness Prevention model showing cost-effectiveness payoffs

In addition, if you double-click on a terminal node to open an Enter Payoff window, the two active payoffs are labeled “Cost” and “Eff.”



Edit cost-effectiveness payoffs

The tree is now ready for cost-effectiveness analysis. The remainder of this chapter will cover performing cost-effectiveness analysis using the TreeAge Pro Healthcare module. The next chapter covers additional cost-effectiveness modeling and analysis options.



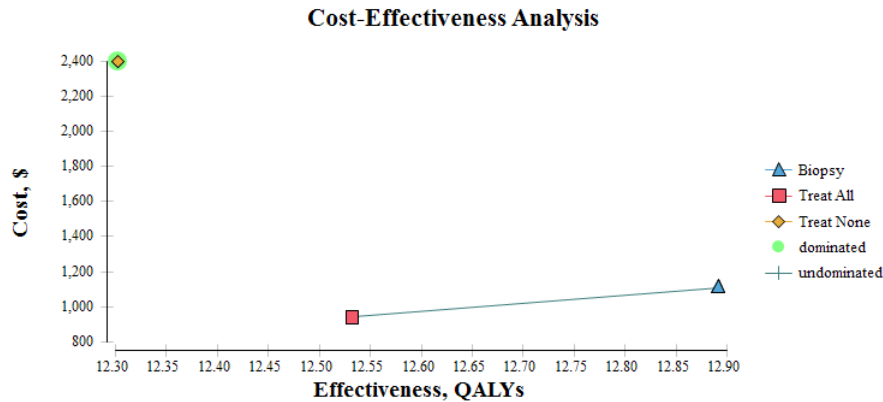
Users of TreeAge Pro who *do not* have the Healthcare module can open trees using the Cost-Effectiveness calculation method, but cannot analyze them. It is possible to change the calculation method of such trees to Simple in order to analyze one of the payoffs, even without the Healthcare module.

31.3 Performing cost-effectiveness analysis

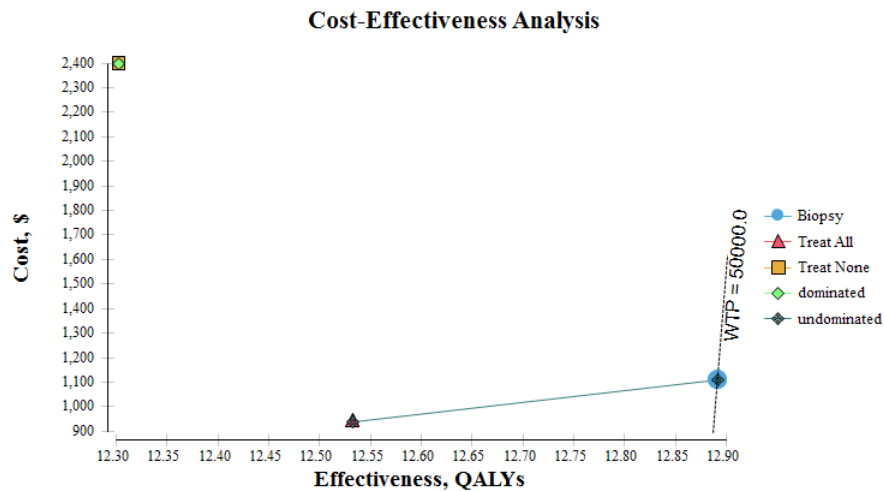
TreeAge Pro’s cost-effectiveness (CE) graphs, and the text report underlying it, are the fundamental tools for cost-effectiveness analysis of your decision trees. They display the key information from the analysis, including incremental values and conditions of dominance.

To generate a CE graph:

- Select the decision node.
- Choose Analysis > Cost-Effectiveness... from the menu.

**Cost-Effectiveness Graph**

The optimal strategy from the cost-effectiveness frontier can be identified using the "Add WTP to Graph" link to the right of the graph. See below.

**Cost-Effectiveness Graph with WTP Line**

The process of interpreting the results of the cost-effectiveness analysis graphically is described in detail later in this chapter. A quick overview is given here.

Strategies which are *not dominated* will be connected by line segments which form the *cost-effective frontier*. Only the strategies on the cost-effective frontier could be the optimal choice. The lowest cost option is always part of this frontier; if it dominates all comparators, the graph will have no lines.

Another option provided is the WTP; this shows the willingness-to-pay slope that intersects with the favored strategy. This can be helpful when it comes to choosing a final strategy.

Details are provided later in this chapter on the rules used to determine which options are excluded from the cost-effective frontier.



You can change the graph x-y orientation of cost and effectiveness via the Tree Preference category C/E Parameters.

31.3.1 The CE analysis text report

To see the calculated values underlying the CE graph, open the graph's text report.

To display the cost-effectiveness analysis text report:

- Click on the Text Report link to the right of the CE graph.

The text report for the Blindness Prevention decision is displayed below.

Category	Strategy	Cost	Incr Cost	Eff	Incr eff	Incr C/E (ICER)	NMB	C/E
▲ Excluding dominated								
undominated	Treat All	940		12.53			375,030	75.01
undominated	Biopsy	1,110	170	12.89	0.36	473.94	385,621	86.11
▲ All								
undominated	Treat All	940		12.53			375,030	75.01
undominated	Biopsy	1,110	170	12.89	0.36	473.94	385,621	86.11
abs. dominated	Treat None	2,400	1,290	12.30	-0.59	-2,192.14	366,677	195.08
▲ All referencing common baseline								
undominated	Treat All	940		12.53			375,030	75.01
undominated	Biopsy	1,110	170	12.89	0.36	473.94	385,621	86.11
abs. dominated	Treat None	2,400	1,460	12.30	-0.23	-6,354.15	366,677	195.08
▲ All by Increasing effectiveness								
abs. dominated	Treat None	2,400		12.30			366,677	195.08
undominated	Treat All	940		12.53			375,030	75.01
undominated	Biopsy	1,110		12.89			385,621	86.11

Cost-Effectiveness Text Report

The Text Report is divided into four collapsible groupings. The columns show a standard cost-effectiveness analysis table, showing average and incremental cost and effectiveness values, as well as incremental cost-effectiveness ratios. This report is described in more detail in the next section.



The Net Monetary Benefits (NMB) is calculated using the WTP parameter from the Tree Preferences. The optimal strategy will have the largest NMB value given that WTP value.



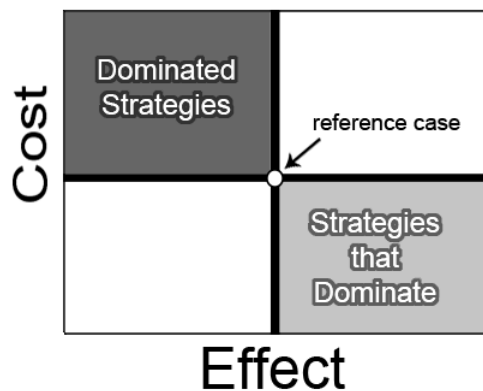
A text report, when generated from a graph window, uses the formatting of the graph axes. To change the formatting of the CE text report, first change the formatting of the graph, then regenerate the Text Report.

31.4 Dominance and incremental cost-effectiveness

In a cost-effectiveness analysis, sometimes a strategy can be eliminated based on its relative cost and effectiveness compared to another strategy. An option is said to be *dominated* if it both costs more

and is less effective than a comparator. This condition can be visually identified in a cost-effectiveness graph.

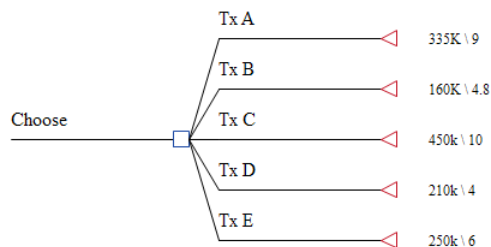
When effectiveness is plotted on the X-axis, a strategy is absolutely dominated if it lies above and to the left of another alternative. The option below and to the right is referred to as dominant, or dominating. TreeAge Pro shows dominated strategies in the graph, but they will not be displayed on the cost-effectiveness frontier.



Dominance

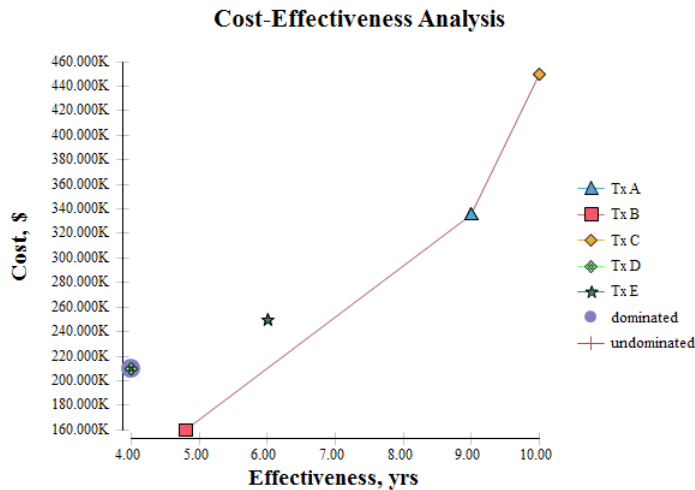
In the CE graph shown in the previous section, the baseline analysis of the Blindness Prevention treatment decision, it is visually apparent that Treat None is dominated by Treat All. (However, we will see later that this may not be the case for all estimates of the uncertain or variable parameters, such as the prevalence of the disease, i.e., if a different population is modeled.)

The Health Care tutorial example model, Extended Dominance, represents the cost and effectiveness of five strategies. The tree has been set up to illustrate special dominance conditions.



Extended Dominance Tree

The CE graph from this tree is shown below.



Extended Dominance CE Graph

By default, extended dominance lines are not shown, however, there is an option to show them.

Tx D is the only option removed from the cost-effective frontier based on absolute dominance — it is more costly and less effective than *Tx B*, the least costly option. The cost-effective frontier is defined starting with *Tx B*, skipping *Tx E*, continuing to *Tx A*, and then to *Tx C* (the most costly and most effective option).

Why is strategy *Tx E* excluded? It is not dominated in the absolute sense — visually, there is no option below and to the right. TreeAge Pro has, instead, flagged *Tx E* as being extendedly dominated, marking it in the graph area (with horizontal and vertical lines intersecting the cost-effective frontier which are hidden in the above graph). Interpreting special conditions of extended dominance in a CEA requires an understanding of the related concept of incremental cost-effectiveness ratios (ICERs).



Extended dominance lines can be hidden by clicking the "Hide Ext. Dom." link to the right of the graph. You can reshown the lines by clicking the "Show Ext. Dom." link.

31.4.1 Extended dominance and ICERs

A single cost-effectiveness analysis for a particular health condition takes place within a wider context; providing the best range of treatments and prevention is the goal, but financial, human, and other resources eliminate some options.

In CEA, when comparing two non-dominated options, an incremental cost-effectiveness ratio (ICER) is calculated. The ICER of the more effective option is the ratio of the mean incremental cost and mean incremental effectiveness (e.g., in terms of \$/QALY). Graphically, it is the slope of the line connecting two, cost-ordered strategies.

$$ICER = IC/IE$$

ICERs are used in the CEA process in a couple of ways. First, ICERs are used to determine whether options can be removed from the cost-effective frontier based on extended dominance. Lower ICERs correspond to better value (i.e., lower cost per unit of additional effectiveness). As shown in the partial graph, *Tx A* is more effective than *Tx E* and has a lower ICER (slope decrease). Thus, the cost-effective frontier connects *Tx B* and *Tx A*, but skips *Tx E*, based on extended dominance.

With effectiveness on the horizontal axis, the slope of the line segment connecting two options corresponds to the ICER. Slopes approaching a horizontal orientation correspond to better (lower) ICERs. In graphs on the previous page, it can be seen that *Tx A* is more effective than *Tx E*, and also has a lower ICER than *Tx E*. In other words, *Tx A* is a better value than *Tx E*, relative to *Tx B*.

In addition to the visual indication of extended dominance provided in the CE graph, the text report provides details about conditions of extended dominance.

Category	Strategy	Cost	Incr Cost	Eff	Incr eff	Incr C/E (ICER)	NMB	C/E
▲ Excluding dominated								
undominated	Tx B	160,000		4.80			80,000	33,333
undominated	Tx A	335,000	175,000	9.00	4.20	41,667	115,000	37,222
undominated	Tx C	450,000	115,000	10.00	1.00	115,000	50,000	45,000
▲ All								
undominated	Tx B	160,000		4.80			80,000	33,333
abs. dominated	Tx D	210,000	50,000	4.00	-0.80	-62,500	-10,000	52,500
ext. dominated	Tx E	250,000	90,000	6.00	1.20	75,000	50,000	41,667
undominated	Tx A	335,000	85,000	9.00	3.00	28,333	115,000	37,222
undominated	Tx C	450,000	115,000	10.00	1.00	115,000	50,000	45,000
▲ All referencing common baseline								
undominated	Tx B	160,000		4.80			80,000	33,333
abs. dominated	Tx D	210,000	50,000	4.00	-0.80	-62,500	-10,000	52,500
ext. dominated	Tx E	250,000	90,000	6.00	1.20	75,000	50,000	41,667
undominated	Tx A	335,000	175,000	9.00	4.20	41,667	115,000	37,222
undominated	Tx C	450,000	290,000	10.00	5.20	55,769	50,000	45,000
▲ All by Increasing effectiveness								
abs. dominated	Tx D	210,000		4.00			-10,000	52,500
undominated	Tx B	160,000		4.80			80,000	33,333
ext. dominated	Tx E	250,000		6.00			50,000	41,667
undominated	Tx A	335,000		9.00			115,000	37,222
undominated	Tx C	450,000		10.00			50,000	45,000

Extended Dominance CE Text Report

Let's look at the CE Text Report in more detail. The collapsible groups are listed below.

1. *Without dominated options*: Only undominated strategies.
2. *All options*: All strategies ordered from least costly to most costly.
3. *All options referenced by a common baseline*: All strategies with incremental values calculated against the common baseline (least costly option).
4. *Ordered by increasing effectiveness*: All strategies ordered from least effective to most effective.

CE Text Report Groups

The columns in the Text report include:

1. *Category*: Grouping from the prior list and whether a strategy is dominated (and if so, how).

2. *Strategy*: The node label for the strategy (branch of the decision node).
3. *Cost*: Cost value for the strategy.
4. *IncrCost*: Incremental cost - difference in cost between this strategy and the previous less costly strategy on the cost-effectiveness frontier.
5. *Eff*: Effectiveness value for the strategy.
6. *IncrEff*: Incremental effectiveness - difference in effectiveness between this strategy and the previous less costly strategy on the cost-effectiveness frontier.
7. *Incr CE (ICER)*: The incremental cost-effectiveness ratio comparing this strategy to the previous less costly strategy on the cost-effectiveness frontier.
8. *NMB*: Net monetary benefits as calculated from the cost and effectiveness for the strategy and the willingness-to-pay (WTP) for the tree. The highest value indicates the favored strategy for the given WTP. More information about the NMB can be found in the next chapter.
9. *Avg CE*: The average cost-effectiveness (Cost divided by Effectiveness) for the strategy.

CE Text Report Columns

ICERs are calculated using only the strategies on the cost-effectiveness frontier, excluding all dominated strategies. Therefore, we will focus on the *undominated* group. The undominated strategies are sorted from least costly to most costly: *Tx B*, *Tx A*, *Tx C*. Incremental values are then calculated for each strategy as compared to the previous less costly strategy.

For *Tx A* compared to *Tx B*...

$$ICER = IC/IE = (335K - 160K)/(9 - 4.8) = (175K)/(4.2) = 41,667$$

For *Tx C* compared to *Tx A*...

$$ICER = IC/IE = (450K - 335K)/(10 - 9) = (115K)/(1) = 115K$$

Note that the ICER, IC and IE values are presented in the Text Report.

In the undominated group, the ICERs will always increase as you move to more costly options. When the ICER decreases in the *all* group, then this indicates that the previous strategy is dominated via *extended dominance*.

Note that the ICER for *Tx E* is larger than the ICER for *Tx A*, which eliminates *Tx E* from the cost-effectiveness frontier due to extended dominance. This indicates that it would be a more efficient use of resources to switch from *Tx B* to *Tx A* than it would be to switch from *Tx B* to *Tx E*.

31.4.2 The threshold ICER (or willingness-to-pay, or ceiling ratio)

The next way in which the ICER is used is to determine if, at some point on the cost-effective frontier, the next more effective option exceeds a threshold ratio, sometimes referred to as the willingness-to-pay, or ceiling ratio. To efficiently allocate resources among competing priorities, there is normally a limit to the additional cost per unit of effectiveness gained – i.e., the ICER – that a rational decision maker will accept to move up the frontier.

If the decision maker assumed a threshold ICER of 40,000, for example, then *tx A* would exceed this — it might be considered slightly inefficient (too costly per unit of effectiveness gained).

The Cost-Effectiveness Modeling and Analysis Options Chapter continues the discussion of CEA thresholds in the context of sensitivity analysis in TreeAge Pro.

31.4.3 Extended dominance: an additional perspective

In some cases, strategy selection may involve not just maximizing effectiveness and working within a threshold ICER, but also working under a budget constraint (i.e., a cost threshold). If such a cost ceiling were set at \$300K, this would eliminate *Tx A* because the cost exceeds the cost ceiling.

Theoretically, if a decision maker is making a population-wide policy decision, two (or more) strategies might be combined to create a “blended” strategy that is less expensive (and less effective) than the too costly option. For example, instead of assigning *Tx A* to all patients, they could be randomly assigned in some proportion to *Tx A* and *Tx B*. In the CE graph, this would result in a new strategy somewhere on the line connecting the two strategies.

The line connecting two options in the graph represents the average cost and effect for all possible blends of the two treatments. The intersection of the ICER line with the cost ceiling (a horizontal line) represents the best hypothetical option blending of *Tx A* and *Tx B*. This optimal blend point is represented as *k* calculated as the ratio:

$$(CostA - CostCeiling)/(CostA - CostB)$$

The interpretation is that *k*% of patients treated are given the less effective Treatment B instead of *Tx A* (or all patients are given *Tx B* for *k*% of their treatments, and *Tx A* the rest of the time). However, questions of equity mean that blends are not often relevant.

The other aspect of the concept of blending, and the blend line, is related to the extended dominance concepts discussed in the previous section. If a blended strategy is created, it may cause an extendedly dominated strategy to become an absolutely dominated one. In the extended dominance example on previous pages, some hypothetical blends of *Tx A* and *Tx B* would absolutely dominate *Tx E*.

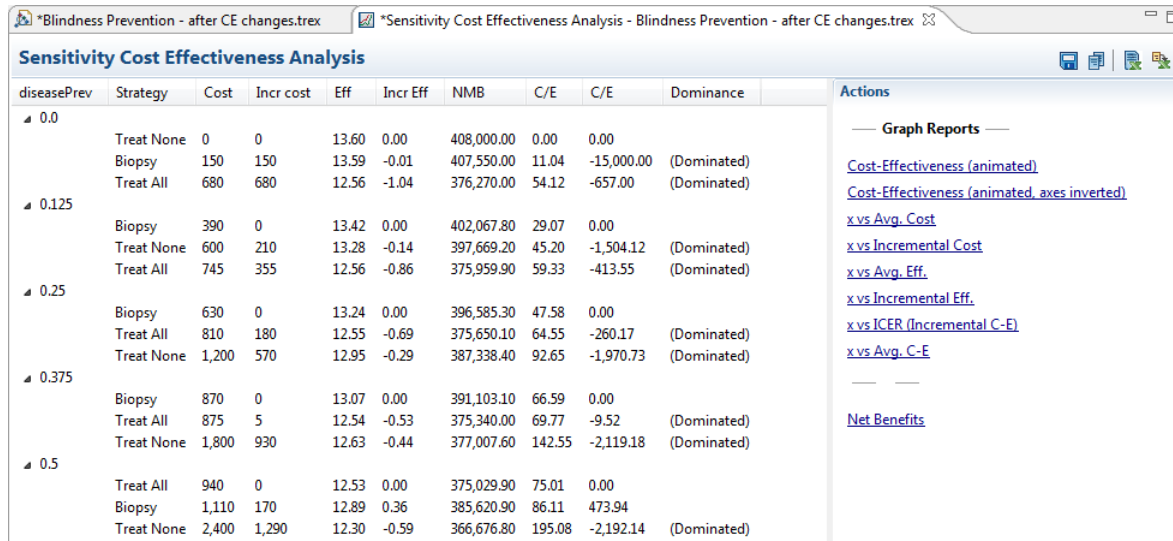
31.5 One-way cost-effectiveness sensitivity analysis

In a CE tree, the steps described in the Introduction to Variables and Sensitivity Analysis Chapter can be used to perform a one-way sensitivity analysis.

Because of the added complexity of multiple attributes and incremental calculations, TreeAge Pro provides several ways to view textual and graphical results of one-way CE sensitivity analysis.

31.5.1 CE sensitivity analysis text report

The immediate output from a CE sensitivity analysis is a text report.



diseasePrev	Strategy	Cost	Incr cost	Eff	Incr Eff	NMB	C/E	C/E	Dominance
0.0	Treat None	0	0	13.60	0.00	408,000.00	0.00	0.00	
	Biopsy	150	150	13.59	-0.01	407,550.00	11.04	-15,000.00	(Dominated)
	Treat All	680	680	12.56	-1.04	376,270.00	54.12	-657.00	(Dominated)
0.125	Biopsy	390	0	13.42	0.00	402,067.80	29.07	0.00	
	Treat None	600	210	13.28	-0.14	397,669.20	45.20	-1,504.12	(Dominated)
	Treat All	745	355	12.56	-0.86	375,959.90	59.33	-413.55	(Dominated)
0.25	Biopsy	630	0	13.24	0.00	396,585.30	47.58	0.00	
	Treat All	810	180	12.55	-0.69	375,650.10	64.55	-260.17	(Dominated)
	Treat None	1,200	570	12.95	-0.29	387,338.40	92.65	-1,970.73	(Dominated)
0.375	Biopsy	870	0	13.07	0.00	391,103.10	66.59	0.00	
	Treat All	875	5	12.54	-0.53	375,340.00	69.77	-9.52	(Dominated)
	Treat None	1,800	930	12.63	-0.44	377,007.60	142.55	-2,119.18	(Dominated)
0.5	Treat All	940	0	12.53	0.00	375,029.90	75.01	0.00	
	Biopsy	1,110	170	12.89	0.36	385,620.90	86.11	473.94	
	Treat None	2,400	1,290	12.30	-0.59	366,676.80	195.08	-2,192.14	(Dominated)

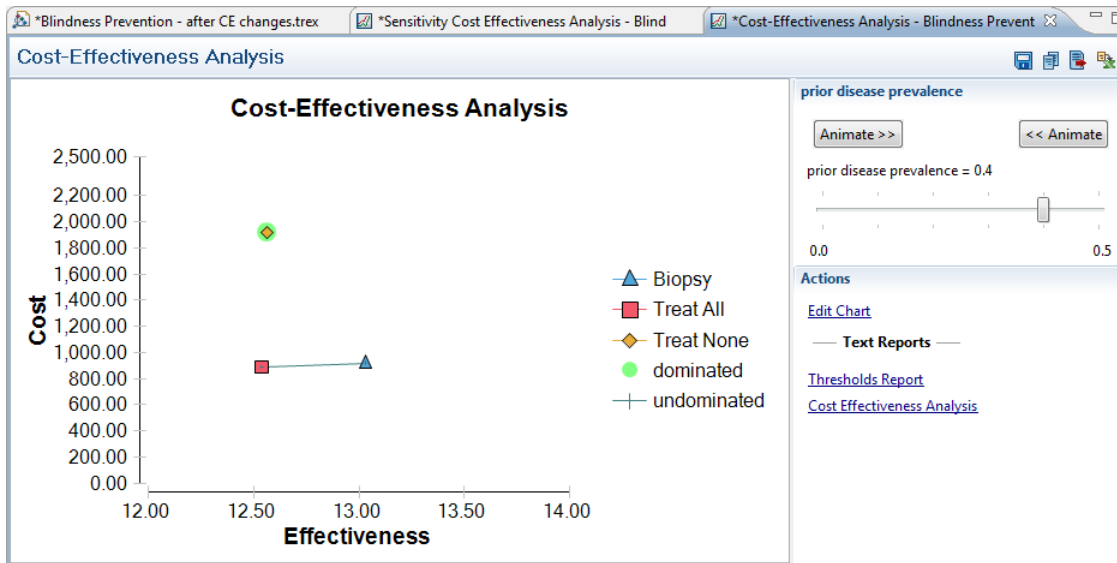
One-Way CE Sensitivity Analysis Output

This text report essentially mirrors the output from the cost-effectiveness analysis text report's *all* group, except with a separate group showing separate CE analysis results for each variable value specified by the sensitivity analysis range/intervals. The NMB column identifies the optimal strategy for that parameter value based on the WTP entered in Tree Preferences. The Dominance column identifies whether a strategy is dominated at that variable value or not.

To the right of the text report are links to generate graphs based on the CE sensitivity analysis data. Several of these will be described in the next section.

31.5.2 CE sensitivity analysis graphs

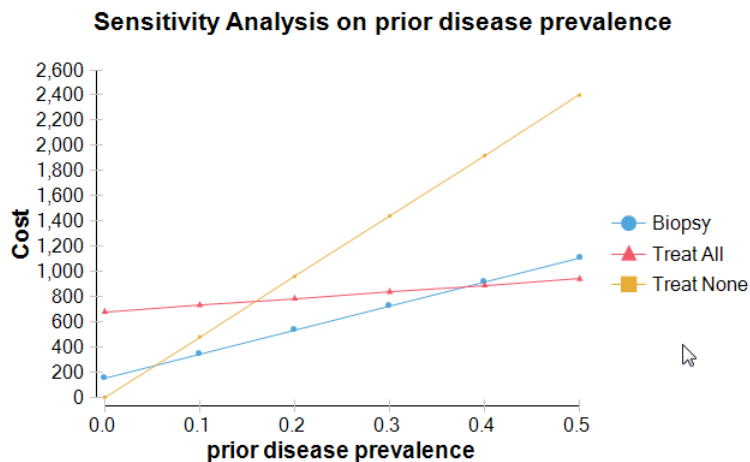
The *Cost-Effectiveness (animated)* link and its inverted version show the cost-effectiveness analysis graph for every value of the variable.



Cost-Effectiveness (animated) graph

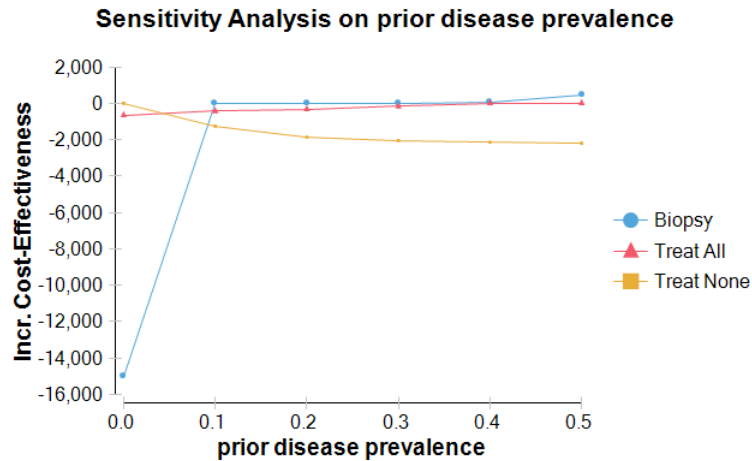
Each frame shows the CE graph for one iteration of the sensitivity analysis. Pressing the Animate buttons, or using the slider, causes TreeAge Pro to step through each value of the sensitivity analysis variable.

Each of the *x* vs "Value" links show how a calculated value changes as the variable changes. The *x* vs Avg. Cost graph below shows how the cost of each strategy changes with the change in the variable diseasePrev.



x vs Average Cost graph

The remaining *x* vs. "Value" graphs look similar, but let's look at the *x* vs. ICER graph.



x vs ICER graph

Note that one of the strategies will always have a zero ICER value at each variable value. Negative ICERs for the other strategies indicates absolute dominance.

The Net Benefits graph may be the easiest to interpret, because Net Benefits calculations combine cost, effectiveness and willingness-to-pay into a single value; the strategy with the highest Net Benefits value is the recommended strategy (given a fixed WTP). This graph is very useful for identifying cost-effective thresholds, particularly in models with small differences in effect, changing cost ordering, and/or more than two strategies. The Cost-Effectiveness Modeling and Analysis Chapter provides details on Net Benefits calculations and the CE sensitivity analysis Net Benefits graph.

32. Cost-Effectiveness Modeling and Analysis Options

The previous chapter provided instructions on preparing a tree for cost-effectiveness calculations, performing baseline cost-effectiveness analysis, and running cost-effectiveness sensitivity analysis.

This chapter covers a variety of other useful cost-effectiveness modeling preferences and techniques available in TreeAge Pro Healthcare and TreeAge Pro Suite, as well as additional cost-effectiveness analysis features.

32.1 Net benefits calculations

The calculation of net monetary benefits (NMB) and net health benefits (NHB) is increasingly prevalent in health economic evaluations, either in addition to or sometimes instead of using ICERs.

In essence, the Net Benefits calculations combine cost, effectiveness and willingness-to-pay into a single measurement. The strategy with the highest Net Benefit is the most cost-effective given the fixed willingness-to-pay parameter.



Willingness-to-pay (WTP) represents how much you are willing to pay for an additional unit of effectiveness. In cost-effectiveness analysis, this is compared to the incremental cost-effectiveness ratio (ICER) to determine if a more expensive treatment should be considered cost-effective.

Willingness-to-pay is a weight on effectiveness in Net Benefits calculations.

The net monetary benefit (NMB) of an alternative is calculated using the following formula:

$$NMB = E * WTP - C$$

Where E represents effectiveness, C represents cost, and WTP is the willingness to pay (i.e., the decision maker's threshold ICER).

The net health benefit (NHB) of an alternative is calculated using a similar formula:

$$NHB = E - C/WTP$$

Some advantages of using net benefits:

- Regardless of the number of strategies, the most cost-effective comparator is simply the one with the highest net benefit, given a threshold ICER ("WTP" in TreeAge Pro).
- When trying to describe uncertainty in CE models with small mean differences in effectiveness (or many competing alternatives), net benefit calculations are not unstable as ratios like the ICER can be.
- Net benefits are the basis for acceptability curves and value of information curves.

Advantages of Net Benefits calculations

Additional background on the net benefits framework and analyses can be found in various journal articles, including:

- "Quantifying stochastic uncertainty" Glick, Briggs, and Polsky, *Expert Rev Pharmacoeconomic Out Res* 1(1), 25-36 (2001) [and www.future-drugs.com].
- "Net Health Benefits," Stinnett and Mullahy, *Med Decis Making* 18 (1998) supplement: S68–S80.

32.2 Multi-attribute weighted costs

TreeAge Pro provides access to an unlimited number of payoff/reward sets. In Multi-Attribute calculations (refer to the Tree Calculation Methods and Preferences Chapter for details), this means that all enabled payoffs can be combined using a weighting function — i.e., a set of numeric or variable *weights* corresponding to pay off or Markov reward sets.

Under the Cost-Effectiveness calculation method, TreeAge Pro allows you to use a weighted cost function in the same way; up to all enabled payoff sets (less one for effectiveness) can be combined as the net cost component of CE calculations.

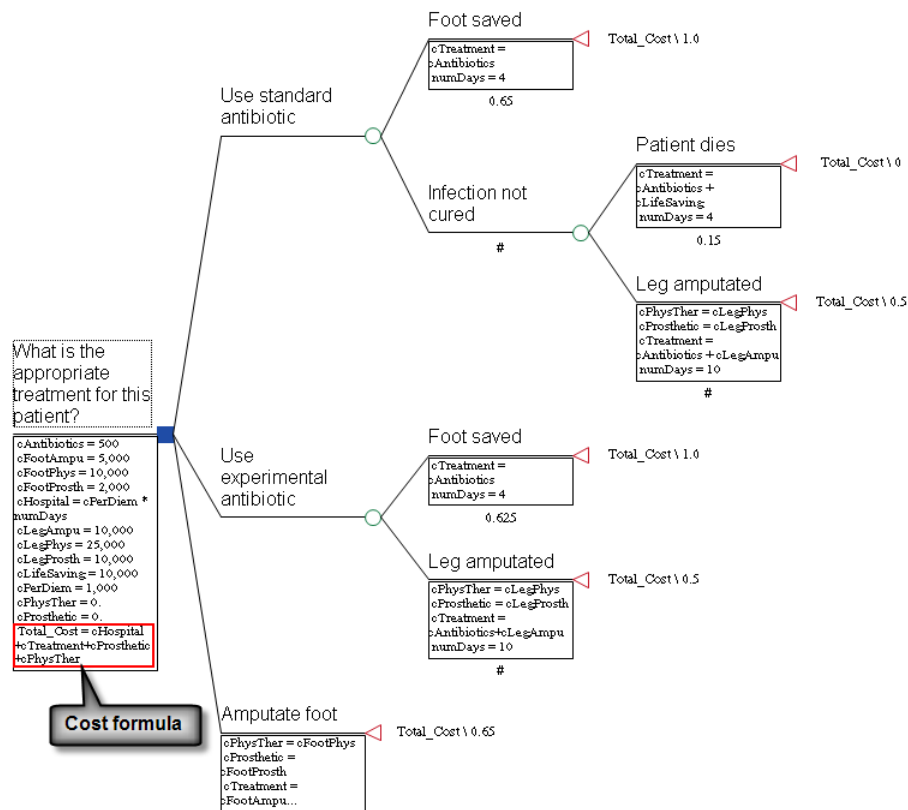
Using a weighted cost function in a cost-effectiveness model may facilitate clearer identification of the parts of a complex cost formula (for example, drug costs, hospital costs, and inpatient costs). It also makes it easier to switch between CEA using a single cost component, and CEA using different combinations of cost components.

32.2.1 Multi-attribute weighted costs: an example

In the example trees in the previous chapter, simple numeric values were assigned to each terminal node's cost payoff. In most models, however, cost payoffs or rewards will be more complicated. The tutorial from the Building Formulas Using Variables and Functions Chapter includes an example tree in which a more realistically complex cost formula is used. The "Cost Formula" tree uses variables to represent the components of a cost formula. In the tutorial, the Simple calculation method was used, but the same issues apply to cost calculations under the Cost-Effectiveness calculation method.

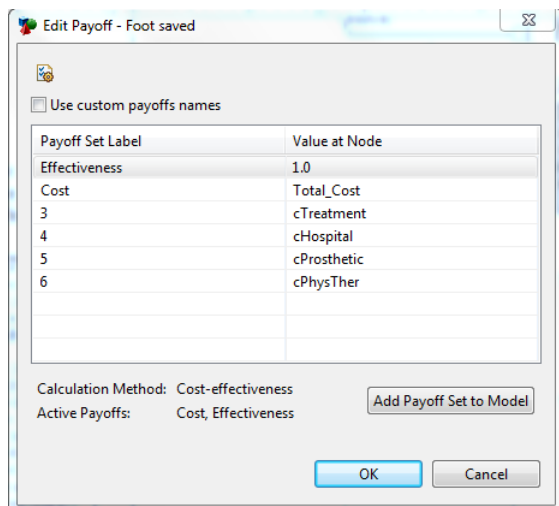
Instead of representing the costs of hospitalization, surgery/drugs, prosthetics, and physical therapy as components of a single payoff (#1 in the example), each of these components can instead be placed in a separate payoff (i.e., #3-#6). Under the Multi-Attribute calculation method or the Cost-Effectiveness calculation method with multi-attribute costs, a simple weighting function could be used to recombine the component variables into a single cost calculation.

To see how the multi-attribute cost weightings work, open the Health Care tutorial example model "CE Cost Formula". This is a CE version of the tutorial example tree "Cost Formula".



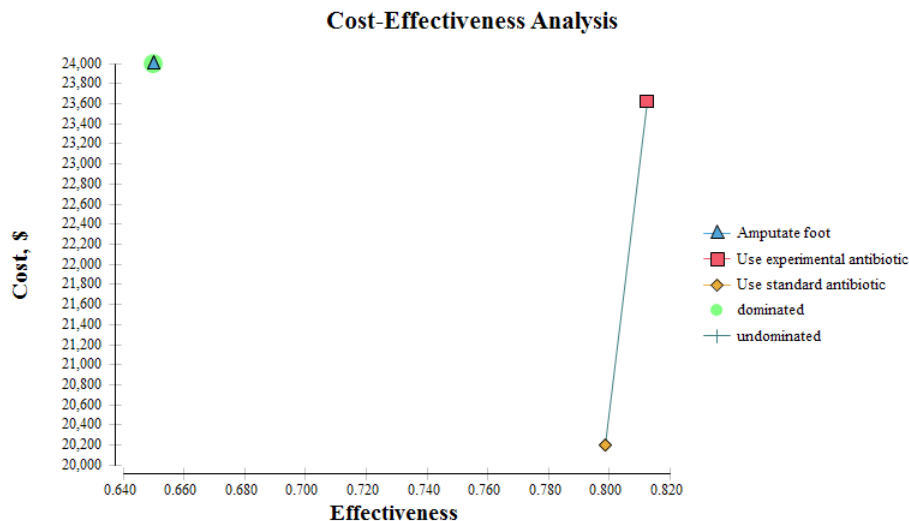
CE Cost Formula tree

If you look at the payoffs for a terminal node, you will see that the current calculation preferences are using the `Total_Cost` variable for cost calculations, which in turn combines several separate measurements of cost. If you change the number of enabled payoffs through the Tree Preferences, it is also possible to see that four other payoffs (#3 through #6) have been assigned the individual components of the `Total_Cost` formula. These separate payoff values can be used for multi-attribute weighted cost calculations.



Terminal Node Multi-Attribute Cost Payoffs

Before making any changes to the tree, perform a cost-effectiveness analysis at the decision node. Later, after changing the payoff calculation preferences in the tree, we can re-run the analysis and compare the results to ensure that no errors were made.

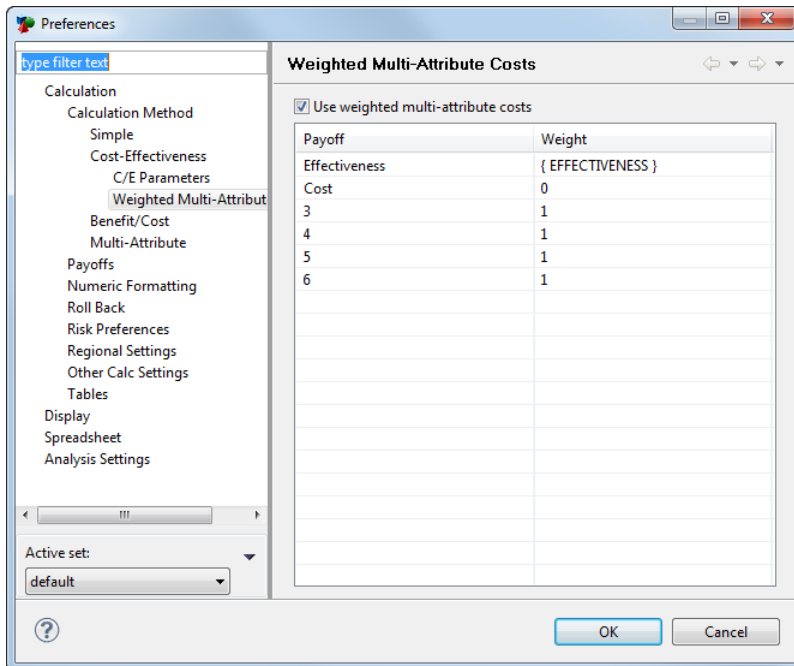


Cost-Effectiveness Analysis on CE Cost Payoff Tree

Now, modify the Calculation Method preferences to use a multi-attribute cost formula, combining the four cost components already in payoffs #3 through #6.

To set up a weighted multi-attribute cost function:

- Choose Tree > Tree Preferences from the menu or press the *F11* key to open the Tree Preferences Dialog.
- Navigate to the category Calculation > Payoffs.
- Note that the number of enabled payoffs to 6.
- Navigate to the category Calculation > Calculation Method.
- Change the Active Method to Multi-Attribute.
- Navigate to the category Calculation > Calculation Method > Cost-Effectiveness > Multi-Attribute.
- Enter a weight of 0 for the Cost payoff and weights of 1 for each payoff that represents a component of the overall cost (payoffs 3-6).
Note that the box corresponding to the effectiveness payoff should be left blank (along with any unneeded payoffs). See below.
- Click OK to save the Tree Preferences.



Tree Preferences - CE Multi-Attribute Costs



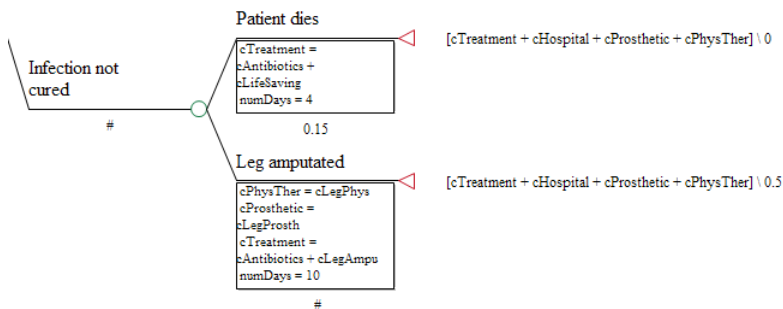
Note that variables could have been entered for each weight instead of 1. This would provide greater flexibility, including the ability to run sensitivity analysis on the weight variables. If using variables for weights, those variables must be defined in the tree; they are then calculated dynamically when analyses are run.

Run another Cost-Effectiveness Analysis to ensure that the results are the same. They should be since the weighted multi-attribute costs mimic the original cost formula for the variable Total_Cost.

Note that a version of this model with the multi-attribute changes is also available - the Health Care tutorial example model: CE Cost Formula - MultiAttribute.trex.

32.2.2 Notes on using multi-attribute costs in CE trees

When weighted, multi-attribute costs are in use in a cost-effectiveness model. Terminal nodes will display the weighted cost payoff formula in brackets; nodes in a Markov model will display separate Markov information line items for each cost reward set.



Multi-attribute cost payoff expressions

To turn off the display of multi-attribute payoff expressions, open the Display > Terminal Nodes preferences category inside Tree Preferences, and uncheck the option labeled Display payoff names.

If you do not enter terminal node payoff expressions for each active cost attribute (i.e., each payoff that is enabled and has an assigned multi-attribute weight), calculation errors will occur. If you enter a weight for the payoff set currently assigned to effectiveness, it is simply ignored during cost calculations. If you leave a weight blank, it evaluates to 0. If you subsequently reduce the number of enabled payoffs in the Calculation Method preferences, any disabled payoffs will be excluded from the weighted cost calculation.

The same weighting function will apply if the calculation method is changed to Multi-Attribute, instead of Cost-Effectiveness.

32.2.3 Markov CE models using multi-attribute costs

Just as with regular trees, Markov models can use the Multi-Attribute calculation method or the Cost-Effectiveness calculation method with the multi-attribute cost preferences described above.

TreeAge Pro calculates each Multi-Attribute (or Cost-Effectiveness with multi-attribute costs) Markov process in a single pass, with the cost weighting done during the Markov process, as well as using either the reward set #1 termination condition (for Multi-Attribute) or the CE termination condition. See the Health Care tutorial example model “Multi Cost Markov”.

See the subsequent chapters on Markov modeling later in the manual for more details.

32.3 Inverting effectiveness calculations

A basic assumption in most CE models is that when it comes to effectiveness, higher numbers are always better. When a cost-effectiveness model presents the reverse situation, with lower values of the effectiveness attribute being preferable, you must invert the calculated effectiveness and/or incremental effectiveness calculations in the tree.

Consider a cost-effectiveness study which tracks the number of adverse events as its measure of effectiveness. In this case, the alternative with the lowest calculated effectiveness value is the most

effective treatment. By default, however, TreeAge Pro's CE analysis algorithm normally identifies options with higher calculated effectiveness value as being preferred. In the case of the adverse event model, leaving this default behavior will result in a CE graph and text report which incorrectly calculates incremental effectiveness and determines dominance.



Net Benefits:

To calculate Net Benefits in trees using inverted incremental effectiveness, TreeAge Pro simply reverses the sign on effects in the Net Benefits calculations. This means that Net Benefits are always negative, but the strategy closest to 0 is optimal.

32.3.1 Inverting incremental effectiveness

When effectiveness measures should be minimized rather than maximized, the strategy selection at decision nodes cannot be done in the standard way. The simplest way to handle this situation in TreeAge Pro is to invert only the incremental effectiveness values for each strategy, so incremental values are inverted.

For example, if a more costly strategy reduces an inverted effectiveness value from 5 to 3, the incremental effectiveness would be calculated as 2 rather than -2. Since the incremental cost is positive, the ICER would also be positive. The positive ICER can then be compared to a WTP threshold.

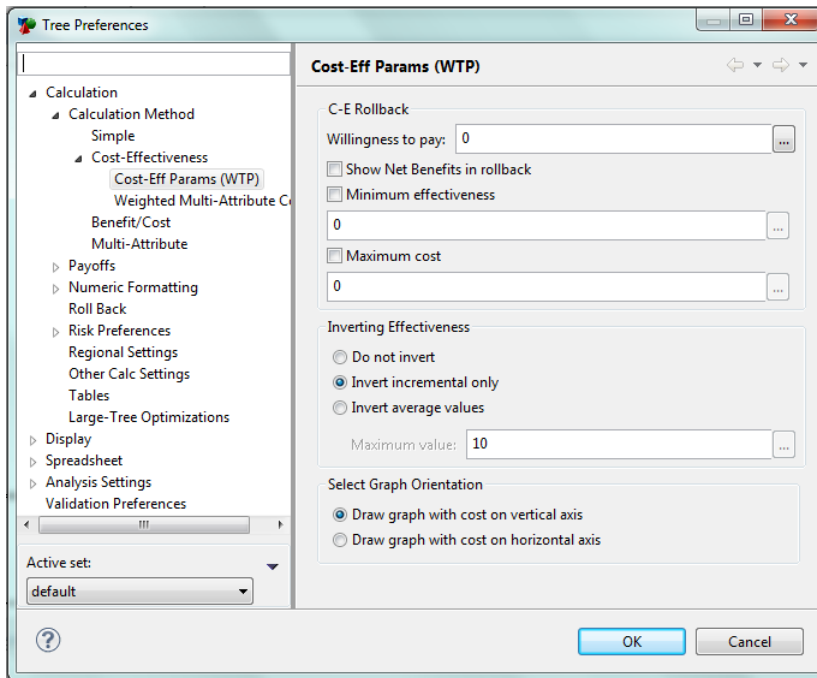
See the Health Care tutorial example model "CE Inverted Effectiveness" for the following steps.

To invert incremental effectiveness calculations only:

- Choose Tree > Tree Preferences from the menu or press the *F11* key to open the Tree Preferences Dialog.
- Navigate to the category Calculation > Calculation Method.
- Confirm that the Cost-effectiveness calculation method is selected.
- Navigate to the category Calculation > Calculation Method > Cost-Effectiveness > Cost-Eff Parameters (WTP).
- Click the option labeled Invert incrementals only.
- Click OK to save the Tree Preferences.

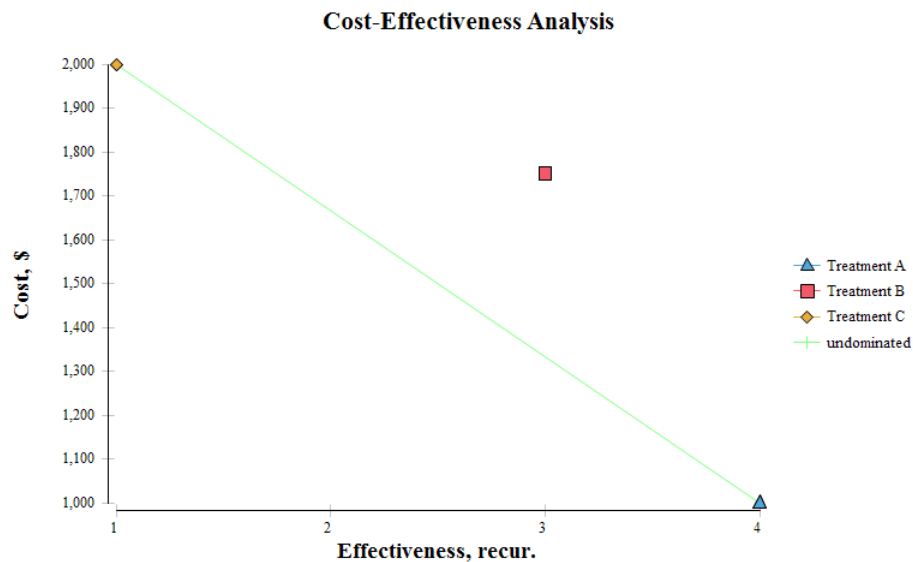


Note that the CE Inverted Effectiveness model already has these preferences set for reference. You can use the steps provided for future work.



Tree Preferences - Cost-Effectiveness - Invert incrementals only

Inverting incremental effectiveness calculations results in a correct construction of the cost-effective frontier (with lines sloping down).



Cost-effectiveness graph with incrementals inverted

Analyses that use or report incremental effectiveness values will simply reverse the normal assumptions, instead of calculating how much "less effective" each more preferable option is.

Cost-Effectiveness Rankings					
Category	Strategy	Cost	Incr Cost	Eff	Incr eff
▲ Excluding dominated					
undominated	Treatment A	1,000		4	
undominated	Treatment C	2,000	1,000	1	3
▲ All					
undominated	Treatment A	1,000		4	
ext. dominated	Treatment B	1,750	750	3	1
undominated	Treatment C	2,000	250	1	2
▲ All referencing common baseline					
undominated	Treatment A	1,000		4	
ext. dominated	Treatment B	1,750	750	3	1
undominated	Treatment C	2,000	1,000	1	3
▲ All by Increasing effectiveness					
undominated	Treatment C	2,000		1	
ext. dominated	Treatment B	1,750		3	
undominated	Treatment A	1,000		4	

Cost-effectiveness text report with incrementals inverted

Inverting incremental effectiveness is usually the preferable method of dealing with an inverted measure of effectiveness; it does not complicate the reporting of expected effectiveness, and does not require specifying a maximum effectiveness value, as does the following, alternate method.

32.3.2 Inverting expected effectiveness calculations

The second method works by inverting the expected effectiveness values calculated for each node, thereby resulting in inverted incremental effectiveness values.

To invert all nodes' effectiveness values:

- Choose Tree > Tree Preferences from the menu or press the *F11* key to open the Tree Preferences Dialog.
- Navigate to the category Calculation > Calculation Method > Cost-Effectiveness > Cost-Eff Parameters (WTP).
- Click the option labeled Invert average values.
- Assign a fixed maximum effectiveness value from which to subtract all nodes' calculated effectiveness values during cost-effectiveness calculations.
- Click OK to save the Tree Preferences.

The inversion of calculated effectiveness values (rather than just incremental values, as above) results, again, in a correct ordering of options in the CE report. The options' calculated incremental effectiveness values (and incremental CE ratios) are the same for both methods.

When inverting all nodes' calculated effectiveness values, a maximum effectiveness value should be selected which is greater than or equal to the uninverted effectiveness of any particular option (even during sensitivity analysis). This will ensure that no inverted values are negative. You may assign a variable or expression for the maximum; at the root node, this expression will be evaluated prior to analysis.

Effectiveness is not inverted *within* a Markov process. TreeAge Pro will invert the Markov node's calculated effectiveness, but Markov analysis graphs and text reports will use *uninverted* effectiveness values.

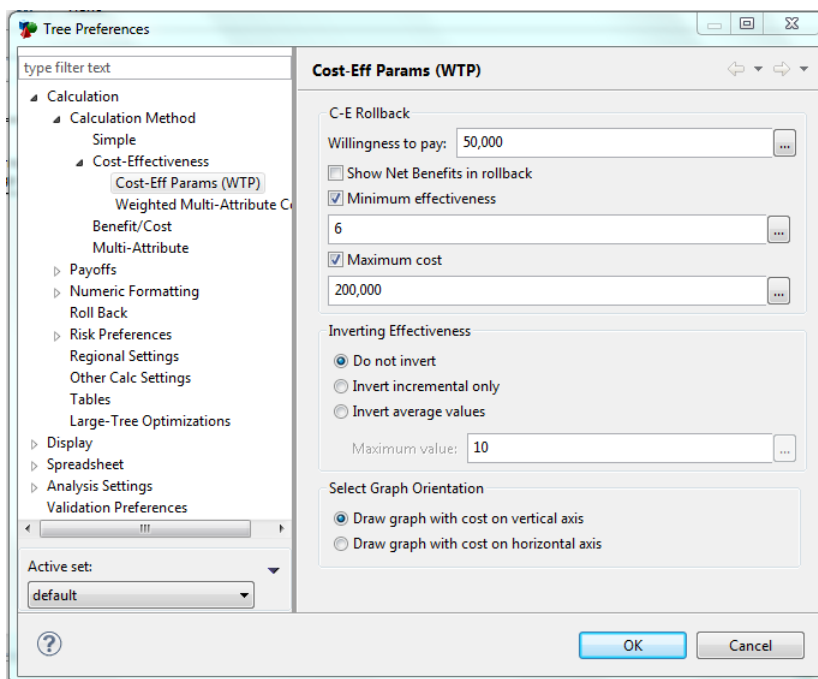
32.4 CE roll back optimal path parameters

As described in the previous chapter, the standard method of performing a baseline CEA in the TreeAge Pro Healthcare module is using the Analysis > Cost-Effectiveness or Analysis > Rankings commands at a decision node. In some cases, however, it may be useful to also display the calculated CEA values in the rolled back tree.

In order to roll back a CE tree, TreeAge Pro must be able to automatically select an optimal path at decision nodes. During CE roll back (and any analysis of a CE tree with embedded decision nodes) TreeAge Pro uses a number of special preferences that enable the model builder to customize the roll back algorithm.

To set the optimal path parameters for cost-effectiveness calculations:

- Choose Tree > Tree Preferences from the menu or press the *F11* key to open the Tree Preferences Dialog.
- Navigate to the category Calculation > Calculation Method > Cost-Effectiveness > Cost-EFF Parameters (WTP).
- Enter values for willingness-to-pay, minimum effectiveness and maximum cost as desired.
- Click OK to save the Tree Preferences.



Tree Preferences - CE Roll Back Parameters

While it is possible to set the CE parameters so that TreeAge Pro makes decisions simply by minimizing the CE ratio, it is also possible to have TreeAge Pro do the following:

- eliminate options below a minimum effectiveness
- eliminate options above a certain cost
- select the most effective option within an incremental cost-effectiveness threshold.

It is also possible to invert the effectiveness measure using the Tree Preferences, as described earlier in this chapter.



Clicking on the ellipses buttons next to the parameter entry boxes will open an expression editor dialog where formulas using variables and functions can readily be set up. Expressions entered for the cost-effectiveness parameters will always be calculated at the root node of the tree, regardless of the location of the decision node being evaluated.

32.4.1 The CE optimal path algorithm

In essence, the optimal alternative will be the most effective option with an ICER lesser than the specified willingness-to-pay (WTP). Any analysis that must select an optimal path, from the strategies at a decision node in a CE tree, does so by using the following algorithm:

1. If minimum effectiveness or maximum cost constraints are specified, any option that fails either test is eliminated.
2. The remaining options are ordered by increasing cost.
3. Any option which is dominated by another less costly, more effective option is eliminated.
4. Each option whose ICER calculated is relative to the next least costly option is greater than your WTP criterion is eliminated.
5. The most effective remaining alternative is selected as optimal.
6. If all options fail these tests, then the least costly option will be selected as optimal.

CE Optimal Path Algorithm



If a WTP of 0 is specified, as is the default, ICERs are ignored, and the least costly option will be selected. If a negative WTP is specified, TreeAge again ignores ICERs, but instead picks the option with the lowest C/E ratio.

32.5 Thresholds and CE sensitivity analysis

The Introduction to Variables and Sensitivity Analysis Chapter covers the use of variables and sensitivity analysis in decision trees. The previous chapter in this manual briefly described the output of a 1-way sensitivity analysis in CE decision trees. This section describes one aspect of CE sensitivity analysis in more detail — finding thresholds. The same Health Care tutorial example model, "Blindness Prevention - after CE changes.trex", is used here.

In a tree, threshold analysis involves searching an uncertain variable's range for values where there is a change in optimal strategy. In a CE model, this means identifying variable values where an alternative changes from being cost-effective to being non-cost-effective (or where the most cost-effective strategy changes).

For a single uncertain variable, CE thresholds can be identified in a number of ways:

- Run a 1-way sensitivity analysis while using either a Net Benefits (NHB or NMB) graph or the Incremental CE graph.
- Run a tornado diagram. Net Benefits calculations are used by default in the series of one-way sensitivity analyses.

Multi-way CE sensitivity analysis can be used to find thresholds when varying 2 or 3 variables simultaneously. By default, Net Benefits are used in dividing graph regions, thus indicating thresholds between regions. As in previous versions of TreeAge Pro, a 2-way analysis on a decision with 2 alternatives can instead be performed with a special ICER isocontour, or threshold, lines (and without Net Benefits calculations).

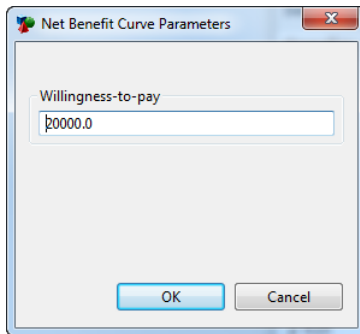
32.5.1 Net Benefits thresholds in a 1-way analysis

Given a particular WTP, a Net Benefits (CE Thresholds) graph is a simple-to-use CE threshold analysis tool. As described at the beginning of the chapter: given the same WTP, the intervention with the highest net benefit (monetary or health) is the most cost-effective. Therefore, thresholds identified by this graph show points within the variable range where there is a change in the most cost-effective strategy.

TreeAge Pro lets you select which scale to use for the vertical axis in the Net Benefits graph, NMB or NHB.

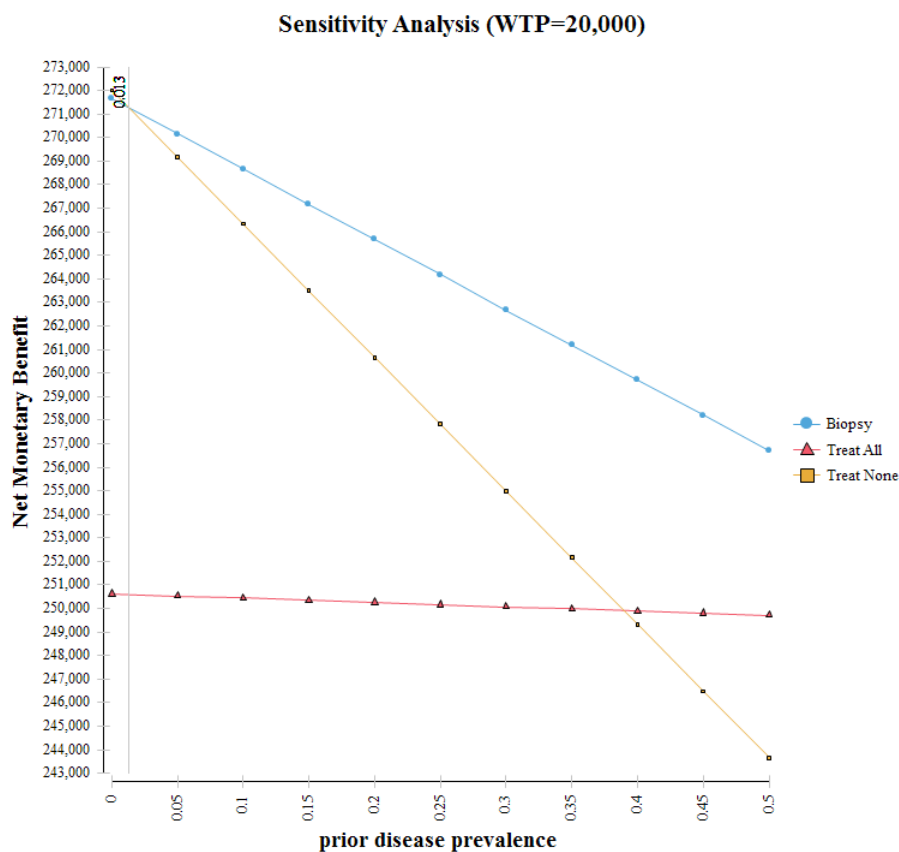
To identify thresholds using the Net Benefits (CE Thresholds) graph:

- Run a one-way CE sensitivity analysis on the Health Care tutorial example model "Blindness Prevention - after CE changes" (variable diseasePrev, range 0-0.5, 10 intervals).
Refer to the prior chapter for details.
- Click the Net Benefits (CE Thresholds) link to the right of the sensitivity analysis output.
- Specify a value for willingness-to-pay (i.e., a ceiling ICER value).



CE Sensitivity Analysis Net Benefits (CE Thresholds) Setup Dialog

A line graph will plot the NMB values for each strategy.



One-way sensitivity analysis Net Benefits (CE Thresholds) graph

If one option's line is always the highest on the vertical benefits scale for a particular analysis, there are no CE thresholds in that case. In this example, however, based on a WTP of 20000, the lines for Treat none and Biopsy exchange places as the option with the highest net benefit at $diseasePrev = 0.013$.

The Thresholds Report (link to right of graph) shows each threshold in the graph in a text format.

1-Way Sensitivity Analysis Thresholds						
Attribute	Variable	Var. Value	Strategy 1	Strategy 2	Exp. Value	Willingness-to-pay
NMB	diseasePrev	0.0116	Treat None	Biopsy	13.56804	30000
Eff	diseasePrev	0.00835	Treat None	Biopsy	13.57832	30000
Cost	diseasePrev	0.05208	Treat None	Biopsy	250	30000
Cost	diseasePrev	0.37857	Biopsy	Treat All	876.85714	30000
NMB	diseasePrev	0.01305	Treat None	Biopsy	13.563	20000

Thresholds Report

Note that thresholds are identified through linear interpolation. Thresholds can usually be identified more accurately by increasing the number of intervals in the original sensitivity analysis.



Several other sensitivity analysis graphs also have a supporting Thresholds Report. They all function in the same way - identifying the points where the maximum (or minimum for cost) value is represented by a change in strategy.

32.5.2 Using a Net Benefits tornado diagram

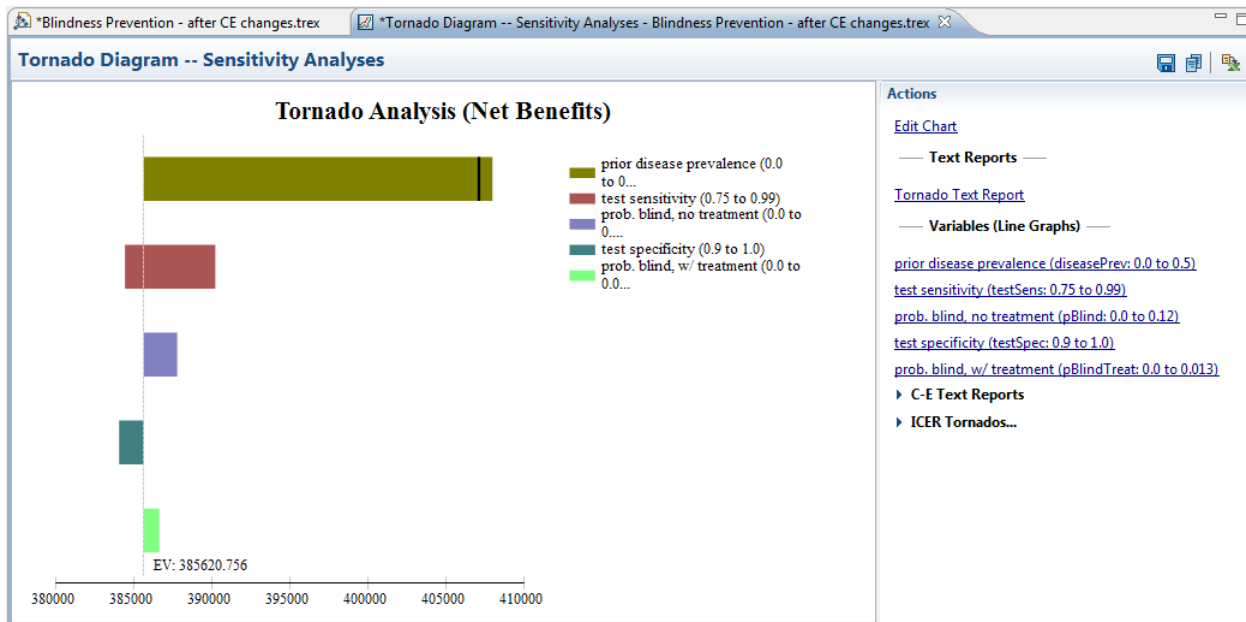
In a tornado diagram on a CE tree, TreeAge Pro normally calculates net benefits (see the beginning of this chapter for details). You can choose which net benefits scale to use for the horizontal axis, monetary or health.

The More Sensitivity Analysis Tools Chapter describes the use of tornado diagrams in detail. This section describes their particular application in CE trees.

To identify thresholds in the Net Benefits tornado diagram:

- Select the decision node.
- Choose Analysis > Sensitivity Analysis > Tornado Diagram from the menu.
- Select the appropriate variable ranges in the setup dialog and click OK.
- Enter a threshold ICER (willingness-to-pay), and select Net monetary benefits or Net health benefits for the y-axis scale.

The following graph was generated from the Health Care tutorial example model "Blindness Prevention - after CE changes".



CE Tornado Diagram

Tornado bars are displayed for each variable, showing how the net benefit of the optimal alternative changes, as well as identifying threshold points with a heavy vertical line. Each bar is rooted on a vertical dotted line indicating the net benefit calculated for the baseline optimal alternative. Click on the Variables (Line Graphs) links to display the underlying one-way sensitivity analysis graph. Clicking on the "prior disease prevalence" link generates the same graph as the one generated in the Net Benefits thresholds in a 1-way analysis section earlier in this chapter.

32.5.3 Creating an incremental cost-effectiveness tornado diagram

After you generate a Tornado Diagram as described in the previous section, you can also create an incremental cost-effectiveness Tornado Diagram via links to the right of the original Tornado Diagram. The ICER Tornado Diagrams report the changing incremental value between two selected strategies (instead of all strategies at a decision node). In a cost-effectiveness sensitivity analysis, this can be used to report incremental cost-effectiveness.

To create an incremental tornado diagram:

- Create a CE Tornado Diagram as described in the previous section.
- Click on one of the ICER Tornado links to the right of the diagram.

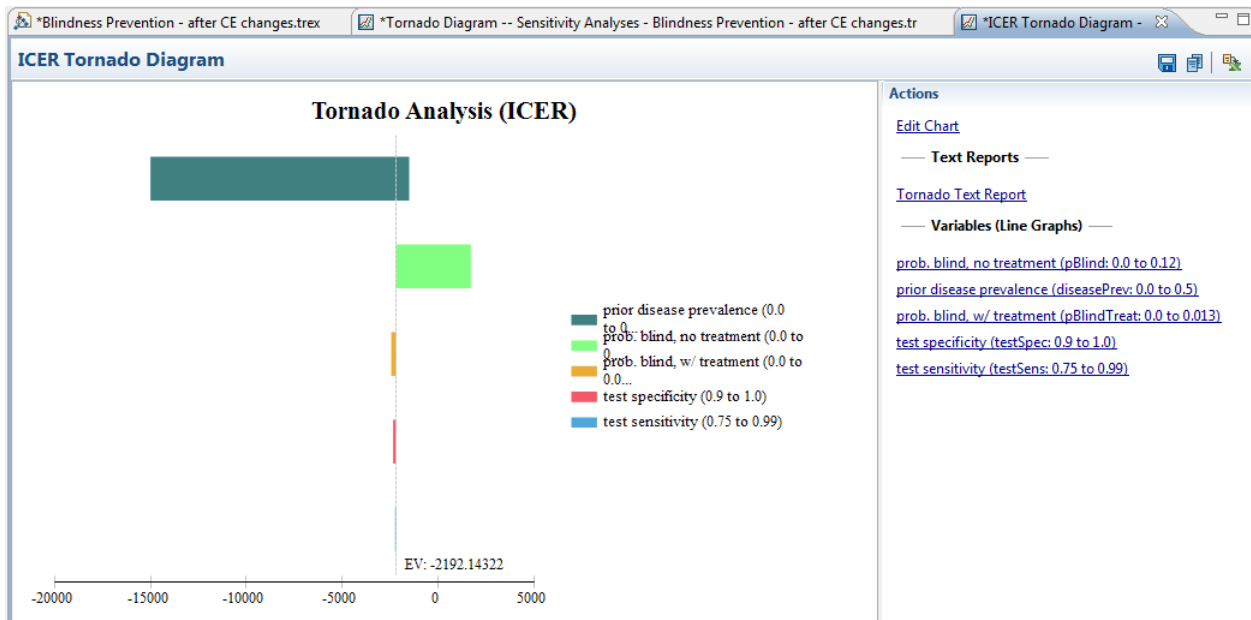
The figure below shows the links generated from the CE Tornado Diagram created in the prior section. Note that there is a separate link for each strategy compared against each of the remaining strategies.

▼ ICER Tornado

[Treat None v. Biopsy](#)
[Treat None v. Treat All](#)
[Biopsy v. Treat None](#)
[Biopsy v. Treat All](#)
[Treat All v. Treat None](#)
[Treat All v. Biopsy](#)

ICER Tornado Diagram Links

The following ICER Tornado Diagram was generated by clicking on the *Biopsy vs. Treat None* link.



ICER Tornado Diagram

As in the original Tornado Diagram, the Variables (Line Graphs) links can be used to show individual one-way sensitivity analysis line graphs - in this case showing ICER vs. the variable values.

32.5.4 Finding cost-effective thresholds using the ICER graph

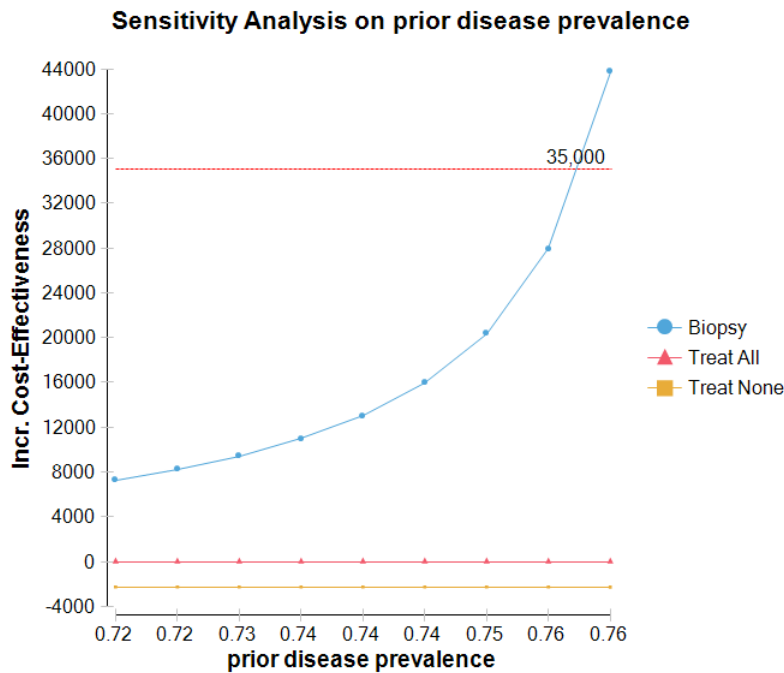
Using TreeAge Pro's incremental CE sensitivity analysis line graph, it is usually possible to find CE thresholds in a more complex way than what was covered in the previous section on net benefits.

The Incremental CE graph shown below is from a one-way sensitivity analysis in the CE version of the Blindness Prevention tree used in the previous chapter.

To generate this graph:

- Open the Health Care tutorial example model "Blindness Prevention - after CE changes.trex".
- Run a 1-way sensitivity analysis on the variable diseasePrev for the range 0.72 to 0.76 with 8 intervals.
- Click the x vs ICER (Incremental C-E) link to the right of the sensitivity analysis output.

After a few modifications (X-axis and added horizontal line marker) as described in the Graph Windows Chapter, the following graph is generated.



Variable vs. ICER Graph

The graph displays the ICERs of three alternatives. For the moment, ignore the two alternatives with horizontal (zero and negative) lines; see below on interpreting zero and negative ICER values in the graph.

The rising curve for Biopsy represents its changing ICER, calculated at each interval relative to the next least costly non-dominated alternative. A dotted horizontal line has been added to the graph to help visualize a WTP (ceiling ratio) of 35,000 \$/QALY. TreeAge Pro will also use the line to approximate the threshold variable value in the Notes section of the text report.

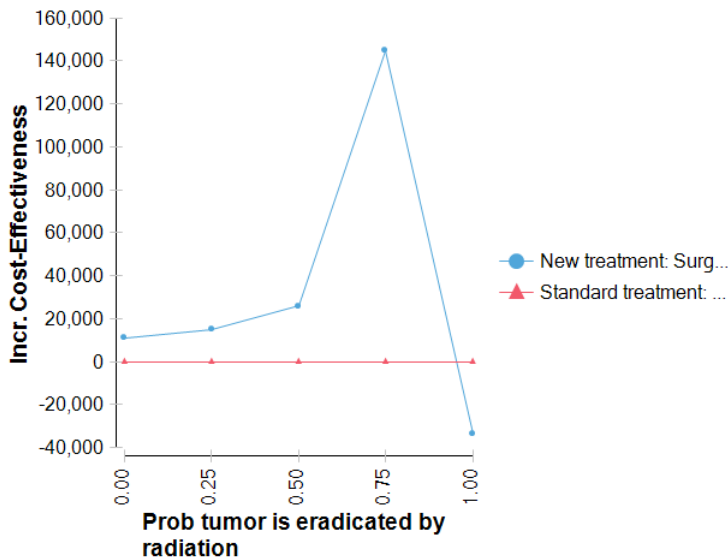
The intersection of the dotted WTP line and the curve representing Biopsy approximates a CE threshold — the prevalence value at which Biopsy changes from being cost-effective to being non-cost-effective, based on a threshold ICER of 35,000, is around 0.76.

Note that the threshold ICER is more easily identified via the Thresholds Report accessible from the Net Benefits sensitivity analysis graph described earlier in this chapter.



Be careful interpreting this graph in cases where the incremental effectiveness approaches zero. This will cause the ICER to approach infinity and negative infinity. Because the graph creates line segments between points based on linear interpolation, the asymptotes in the graph will not be shown accurately.

Sensitivity Analysis on Prob tumor is eradicated by radiation



ICER sensitivity analysis graph with "asymptote"

Note that if you select a range and intervals that calculate a value where IE is exactly equal to zero, the graph will display this value as zero.

The Net Benefits sensitivity analysis graph is not affected by asymptotes, so this is often a better option.

32.5.5 2- and 3-way CE sensitivity analysis thresholds using Net Benefits

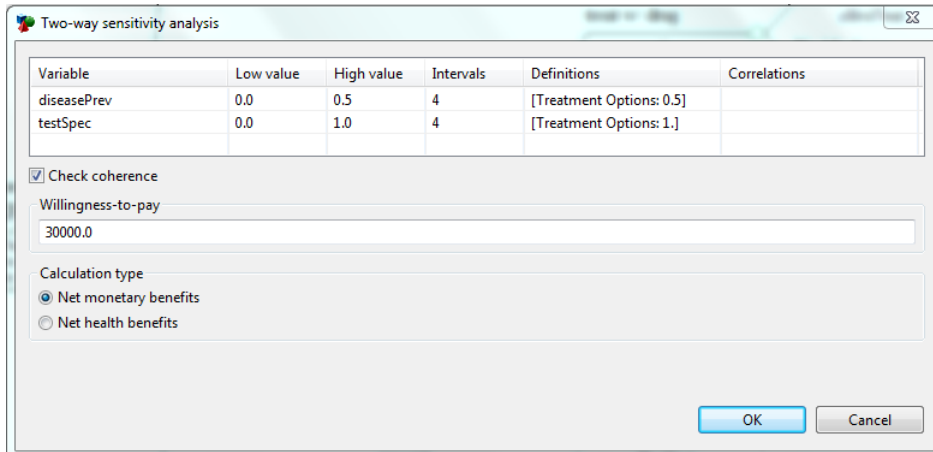
As described in the More Sensitivity Analysis Tools Chapter, in a tree set to calculate a single attribute (e.g., cost or utility), a two-way sensitivity analysis identifies the optimal alternative for each combination of values of the two variables. Based on this, a region graph is created in the two-dimensional variable space with regions assigned to the alternatives based on their optimality. The lines dividing two regions are threshold lines.

It is also possible to run 2- and 3-way sensitivity analysis on cost-effectiveness trees using Net Benefits calculations, making CE thresholds easy to identify.

TreeAge will prompt you for a willingness-to-pay value to use in the Net Benefits calculations for all strategies.

To run 2-way sensitivity analysis on an example model:

- Open the Health Care tutorial example model "Blindness Prevention - after CE changes.trex".
- Select the root node.
- Choose Analysis > Sensitivity Analysis > 2 Way... from the menu.
- Select the variables diseasePrev and testSpec and set the ranges based on the figure below.
- Enter the Willingness-To-Pay parameter and select the calculation type based on the figure below and click OK.



Two-way sensitivity analysis

Variable	Low value	High value	Intervals	Definitions	Correlations
diseasePrev	0.0	0.5	4	[Treatment Options: 0.5]	
testSpec	0.0	1.0	4	[Treatment Options: 1.]	

☒ Check coherence

Willingness-to-pay
30000.0

Calculation type

☒ Net monetary benefits

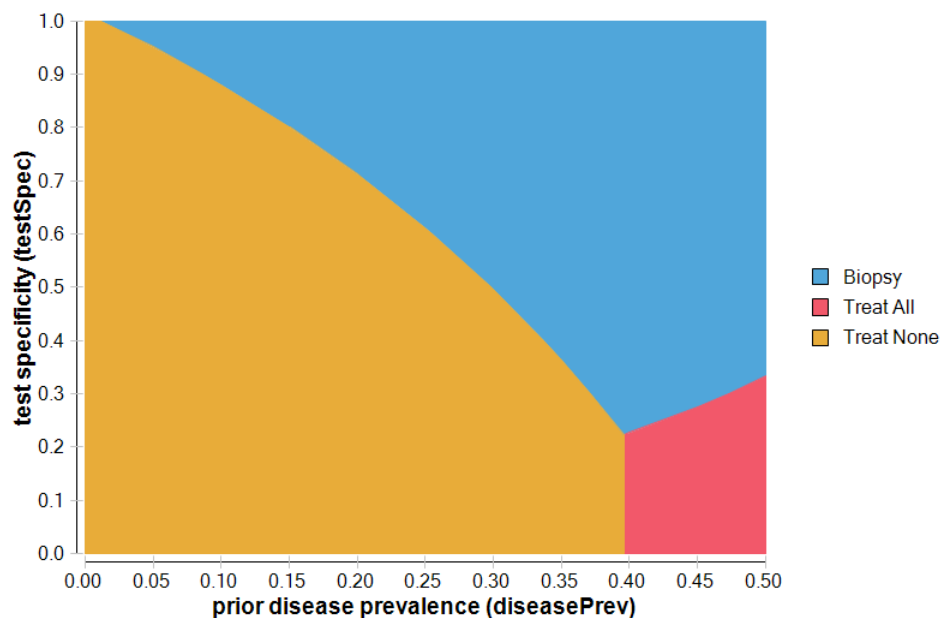
☐ Net health benefits

OK Cancel

2-way sensitivity analysis parameter input dialog

For the given WTP, the strategy having the highest net benefit for any coordinate in the analysis will be assigned that point. Regions of cost-effectiveness can then be constructed on this basis. We entered very large ranges for our two variables, so all three strategy options come into play.

Sensitivity Analysis on diseasePrev and testSpec (Net Benefit, WTP=30000.0)



2-way sensitivity analysis graph

Most of the variable value combinations recommend the *Biopsy* strategy. However, when the test sensitivity is close to zero, the recommended strategy is either *Treat None* or *Treat All*. This is not surprising since a test with no sensitivity has no value.

The Text Report for the graph shows cost, effectiveness and net benefit calculations for each strategy at each combination of variables.

test specificity \ prior disease prevalence	0	0.05	0.1	0.15	0.2	0.25	0.3	0.35	0.4	0.45
0.2										
NMB(30000.0), Treat None	408,000	403,867.68	399,735.36	395,603.039...	391,470.72...	387,338.39...	383,206.08	379,073.7599...	374,941.44	370,809.12
Cost	0	240	480	720	960	1,200	1,440	1,680	1,920	2,160
Eff	13.6	13.470256	13.340512	13.210768	13.081024	12.95128	12.821536	12.691792	12.562048	12.432304
NMB(30000.0), Biopsy	382,166	381,242.2...	380,318.5512...	379,394.8268	378,471.10...	377,547.378	376,623.6536	375,699.9292	374,776.2048	373,852.4804
Cost	694	762.8	831.6	900.4	969.2	1,038	1,106.8	1,175.6	1,244.4	1,313.2
Eff	12.762	12.733502...	12.70500504	12.67650756	12.64801008	12.6195126	12.59101512	12.56251764	12.53402016	12.50552268
NMB(30000.0), Treat All	376,270	376,145.9...	376,021.9860...	375,897.979	375,773.972	375,649.965	375,525.958	375,401.9509...	375,277.944	375,153.937000...
Cost	680	706	732	758	784	810	836	862	888	914
Eff	12.565	12.5617331	12.5584662	12.5551993	12.5519324	12.5486655	12.5453986	12.5421317	12.5388648	12.5355979
0.1										
NMB(30000.0), Treat None	408,000	403,867.68	399,735.36	395,603.039...	391,470.72...	387,338.39...	383,206.08	379,073.7599...	374,941.44	370,809.12
Cost	0	240	480	720	960	1,200	1,440	1,680	1,920	2,160
Eff	13.6	13.470256	13.340512	13.210768	13.081024	12.95128	12.821536	12.691792	12.562048	12.432304
NMB(30000.0), Biopsy	378,993	378,227.9...	377,462.8512...	376,697.776...	375,932.70...	375,167.62...	374,402.5536	373,637.4792	372,872.4047...	372,107.3304
Cost	762	827.4	892.8	958.2	1,023.6	1,089	1,154.4	1,219.8	1,285.2	1,350.6
Eff	12.6585	12.635177...	12.61185504	12.58853256	12.56521008	12.5418876	12.51856512	12.49524264	12.47192016	12.44859768
NMB(30000.0), Treat All	376,270	376,145.9...	376,021.9860...	375,897.979	375,773.972	375,649.965	375,525.958	375,401.9509...	375,277.944	375,153.937000...
Cost	680	706	732	758	784	810	836	862	888	914
Eff	12.565	12.5617331	12.5584662	12.5551993	12.5519324	12.5486655	12.5453986	12.5421317	12.5388648	12.5355979
0										
NMB(30000.0), Treat None	408,000	403,867.68	399,735.36	395,603.039...	391,470.72...	387,338.39...	383,206.08	379,073.7599...	374,941.44	370,809.12
Cost	0	240	480	720	960	1,200	1,440	1,680	1,920	2,160
Eff	13.6	13.470256	13.340512	13.210768	13.081024	12.95128	12.821536	12.691792	12.562048	12.432304
NMB(30000.0), Biopsy	375,820	375,213.5...	374,607.1512	374,000.726...	373,394.30...	372,787.87...	372,181.4536	371,575.0291...	370,968.6048	370,362.180399...

2-way sensitivity analysis text report

Note that at the point where $diseasePrev = 0.4$ and $testSpec = 0.2$, the net benefit values for all three strategies are very close, mirroring the border of all three strategies on the region graph.



Custom isocontours added to the graph will represent incremental net benefits, not incremental CE ratios.

32.5.6 2-way CE sensitivity analysis, cost regions and isocontours

Isocontours have not yet been implemented in TreeAge Pro 201x. This section will be updated when the feature is available.

32.6 Displaying incremental values during roll back

The calculated incremental values of competing strategies in a cost-effectiveness tree are not automatically displayed when the tree is rolled back. The easiest way to report incremental expected values is to generate the CE analysis text report as described in the previous chapter. It is possible, however, to display incremental values in the rolled-back tree.

For a visual display of incremental values in the rolled-back tree, you need to create terminal node columns that display the appropriate incremental values, and then collapse the subtrees to the right of the decision node. Terminal node columns, covered in detail in the Tree Display Preferences and Options Chapter can be used to display incremental values to the right of *visual* end nodes (not just terminal nodes) during roll back.

33. Cost-Effectiveness Simulation Reports and Graphs

The Monte Carlo Simulation Chapter covers the basic aspects of performing Monte Carlo simulation: using the simulation output window to display statistical information, a full text report, and graphs describing the probability distribution of inputs and outputs.

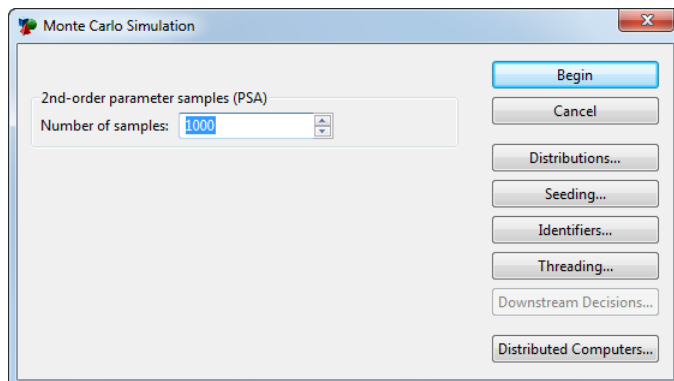
TreeAge Pro Healthcare and TreeAge Pro Suite add a number of graphs and reports designed specifically for Monte Carlo simulation of cost-effectiveness models. These are described in this chapter.

33.1 Basic CE statistics and simulation outputs

As described in the earlier simulation chapter, Monte Carlo simulations generate outputs that present simulation statistics and provides access to a number of reports and graphs. Simulation outputs can be saved for reference later.

To generate CE Monte Carlo simulation output:

- Open the Health Care tutorial example tree "CE Markov Sampling".
- Select the root node.
- From the menu, choose Analysis > Monte Carlo Simulation > Sampling (Probabilistic Sensitivity)...
- Enter 1000 for the number of samples.
- Click Begin.



Simulation setup dialog

Since the simulation was run on a cost-effectiveness tree, statistics for *both cost and effectiveness* are displayed and split up into three groups:

- *Cost*: Cost values.
- *Eff*: Effectiveness values.
- *NMB*: Net Monetary Benefits values.

33.1.1 Simulation output common to simple and cost-effectiveness models

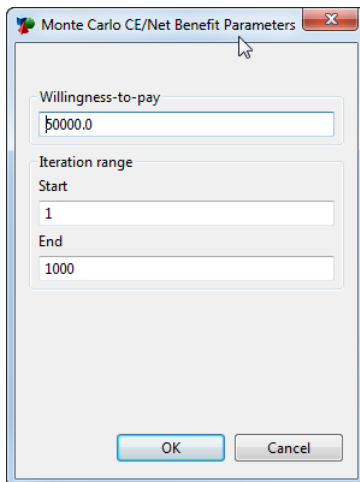
A few output options are described in the Monte Carlo Simulation, Distributions and Probabilistic Sensitivity Analysis Chapter that function exactly the same way for cost-effectiveness models, so they are not described here. These options include:

- Identifying variables
- Outputting the statistics report
- Saving simulation output
- Sampling distributions report and chart output

Several options that appear both for PSA of simple and cost-effectiveness trees are described in this chapter because they function differently within each context.

33.1.2 Simulation text reports

When you click on the *Strategy Values* link, you are prompted for input by the Monte Carlo CE/Net Benefit Parameters dialog.



Monte Carlo CE Parameters Dialog

The Willingness-to-pay and Calculation type parameters control the calculation of Net Benefits information that is presented with each strategy. The Iteration range determines how many of the simulation iterations to present. Given the entries above, the following output is generated.

Monte Carlo Strategy Values

Showing page 1 of 40

Sample	Strategy	Cost	Eff	NMB
1	Rx A	111,123.48	5.31	154,184.42
	Rx B	34,109.24	2.97	114,355.39
	Rx B-2	29,506.65	2.57	98,924.62
2	Rx A	68,357.92	5.55	209,007.58
	Rx B	34,649.47	2.83	106,748.61
	Rx B-2	52,424.87	4.28	161,511.35
3	Rx A	67,169.44	5.79	222,393.16
	Rx B	27,256.86	2.71	108,103.95
	Rx B-2	35,148.64	3.49	139,403.67
4	Rx A	58,793.48	5.74	228,062.99
	Rx B	33,139.14	3.35	134,549.89
	Rx B-2	29,513.20	2.99	119,828.04
5	Rx A	78,613.37	5.54	198,221.68
	Rx B	50,623.79	2.29	63,876.95
	Rx B-2	35,958.00	1.63	45,371.70

Report Options...
[Decimal format](#)
 More Reports
[Values \(html\)](#)

CE PSA Output - Strategy Values

The output shows the cost, effectiveness and net benefits for each simulation iteration in the specified range. The net benefits value is calculated based on the WTP specified in the parameters dialog.

The *Values, Dists, Trackers* link generates a complete set of outputs from the simulation.

Monte Carlo Simulation Report

ITERATION	STRATEGY_1_COST	STRATEGY_1_EFF	STRATEGY_2_COST	STRATEGY_2_EFF	STRATEGY_3_COST	STRATEGY_3_EFF	DIST_1	DIST_2
1	105530.9201020525	5.8132232889	32345.4021806719	2.8054612166	20442.4290623858	1.773063188	17015.1780023499	8780.152458
2	71032.0932993915	5.0033627287	51475.6817220297	3.2200583808	18669.2534207124	1.1678540998	11452.7923209604	13973.06272
3	61031.5643256914	5.4328957004	39587.2842115573	3.3677612247	100494.0719370152	8.5492108268	9840.3664988353	10745.95977
4	59949.2756026192	4.9708043188	32932.2781767136	2.9630828833	77826.8638748624	7.0024748052	9665.8647011139	8939.459820
5	67403.1020828933	5.7634324105	33049.0544362842	2.6515973435	26074.9281426651	2.0920480593	10867.6753575342	8971.158711
6	65012.8688090442	5.6652983848	46115.0940084941	2.7683927855	17285.8104260397	1.0377060679	10482.2883583272	12517.93234
7	80600.7693850012	4.6570539748	27141.680974514	2.809771158	21869.9209540066	2.2640260631	12995.5887514855	7367.603457
8	64030.7273925286	5.234445785	46804.3959943116	3.2559089076	27778.152488376	1.9323640911	10323.9337167745	12705.04321
9	63323.028863863	5.2483507771	35888.7496126361	3.5122298646	44564.69087999	4.3612953893	10209.8286144443	9741.993355
10	73839.1946411321	5.4296165858	39438.2711394561	2.5775204396	41436.624506883	2.708124457	11905.3926484678	10705.51020
11	82776.3491218744	5.7673473784	50146.6911773748	3.1046313228	56281.7702574272	3.4844601457	13346.3662908588	13612.30852
12	82039.9021671988	5.8543478617	46821.4777183959	3.0686064626	42931.3471586773	2.8136533864	13227.6259632755	12709.68004
13	50377.9969873672	5.1232406626	42852.2545584987	2.2697748594	23615.7294620148	1.2508650845	8122.6486541855	11632.23527
14	81025.5607768915	5.5734301607	35092.0314692492	2.9854182774	29719.6866464439	2.5283715989	13064.0795894303	9525.724387
15	65280.289011914	5.0266672009	47150.4369381889	3.1704665006	76151.4650293574	5.1205393741	10525.405601585	12798.97595
16	75991.8604235632	5.6339866397	38488.4244363949	2.8498493403	102836.6437359816	7.6144696905	12252.4756780881	10447.67452
17	92696.3813205854	5.951192092	35964.35446874	3.299455611	48911.2083750478	4.4872308512	14945.8132916704	9762.516277
18	52019.2879829309	5.9107841184	24361.6907021762	2.6250072423	27936.2180527457	3.0101677099	8387.2806541355	6612.975696
19	87151.9770048635	5.7035398398	49311.8404356087	3.4275304936	63150.5437349083	4.3894207238	14051.8665104069	13385.68846
20	80835.3772121629	4.9838713381	53382.7841218756	2.6575937455	81424.7864065416	4.0536290237	13033.4155224076	14490.74525

Strategy Key
 STRATEGY 1 => Rx A
 STRATEGY 2 => Rx B
 STRATEGY 3 => Rx B-2

Distributions Key
 DIST 1 => distribution1
 DIST 2 => distribution2
 DIST 3 => distribution3
 DIST 4 => distribution4
 DIST 5 => distribution5

CE PSA Output - Values, Dists, Trackers

The *Values, Dists, Trackers (Tab-Delimited XLS/TXT)* link creates a report which will export directly to *.csv/*.txt. This is much more efficient for large data output than running the Values, Dists, Trackers report first, then exporting that to Excel.

The output is the same as for simple trees, except that it includes both cost and effectiveness EV values rather than a single EV. The same is true for the *Statistics - Values, Dists, Trackers* link as presented below.

STATISTIC	COST_STRATEGY_1	EFF_STRATEGY_1	NMB_STRATEGY_1	COST_STRATEGY_2	EFF_STRATEGY_2	NMB_STRATEGY_2
Mean	74125.0180064702	5.3822306131	-74125.0180064702	36968.4302418399	2.95379724	-36968.4302418399
Std Deviation	15611.7355450378	0.4002797003	15611.7355450378	7252.1477994248	0.3002662364	7252.1477994248
Minimum	15318.7172015602	4.4058520747	-123109.211540976	16304.3375281471	2.2697748594	-59445.896717801
2.5%	43282.8902318393	4.56553809	-103878.425795661	23294.8915818518	2.3977931775	-51466.6061169517
10%	53985.7274531266	4.7942085069	-93326.7566196121	27477.4292797375	2.5453874523	-46409.888720118
Median	74139.4115089874	5.4317570456	-74318.3747177579	37109.411632498	2.9568009222	-37142.953889989
90%	93319.1010378569	5.8720845461	-54099.0424518648	46309.1097956596	3.3634281175	-27482.8560307129
97.5%	103878.425795661	6.0533017426	-43282.8902318393	51466.6061169517	3.5284911253	-23294.8915818518
Maximum	123109.211540976	6.1753571219	-15318.7172015602	59445.896717801	3.6689503195	-16304.3375281471
Sum (n*Mean)						
Size (n)	1000	1000	1000	1000	1000	1000
Variance	243726286.72819564	0.1602238385	243726286.72819585	52593647.70470249	0.0901598127	52593647.704702474
Variance of Mean (var/n)	243726.2867281956	0.0001602238	243726.2867281958	52593.6477047025	0.0000901598	52593.6477047025
Std Error of Mean	493.6864255053	0.0126579555	493.6864255053	229.3330497436	0.0094952521	229.3330497436

CE PSA Output - Statistics - Values, Dists, Trackers

The interpretation of each value in the simulation *Values, Dists, Trackers* output will depend on the type of simulation you have performed. TreeAge Pro includes a variety of Monte Carlo simulations: one-, two-, or three-dimensional; sampling; microsimulation; and other combinations of these. The Monte Carlo simulation output will report a summary for the “highest” dimension.

In the simplest case, such as a one-dimensional simulation, TreeAge Pro will perform a series of recalculations of the model by either *expected value recalculations* (second-order sampling, PSA) or *individual microsimulations* (first-order trials), and report each result. On the other hand, in a two-dimensional simulation, each row of values reported by TreeAge Pro is itself a summary of a group of recalculations of the model (again, either EV calculations or microsimulation trials). This section focuses primarily on a CE probabilistic sensitivity analysis.

Note that neither of the last two outputs includes tracker data because the model is not a Microsimulation model with trackers. A microsimulation model would include tracker data.

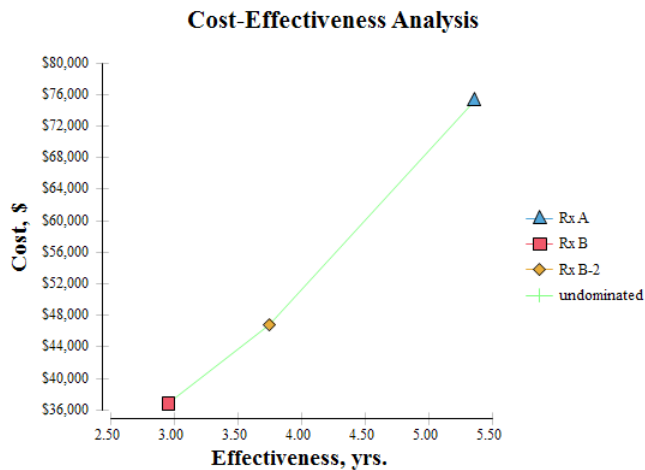
33.2 Cost-effectiveness graphs

CE simulations provides three cost-effectiveness graphs, accessible under the Plots/Curves > CE Analysis grouping.

- *CE graph*: Standard cost-effectiveness graph.
- *CE Graph (inverted)*: Same as above but with axes inverted.
- *CE Rankings Report*: CE text report.

The Monte Carlo simulation *CE Graph* is the same graph as is generated by the Analysis > Cost-Effectiveness menu command. In the simulation version, each strategy is plotted using the mean cost

and effectiveness statistics from the simulation summary, rather than using the expected values. The graph includes the standard CEA graph and report format.



CE PSA Output - CE Graph

See the previous two chapters in this manual for detailed descriptions of the cost-effectiveness graph and text report.

The *CE Rankings Report* option generates a ranked version of the CEA text report.

Cost-Effectiveness Rankings										
Subset	Rank	Strategy	Cost	Incr Cost	Eff	Incr Eff	Incr CE (ICER)	Avg CE	Dominance	Orig. Order
all	1	Rx B	36968.4302418399	0	2.95379724	0	0	0		1
	2	Rx B-2	46510.1635413636	9541.7332995238	3.7179836328	0.7641863928	12486.1334747069	0		2
	3	Rx A	74125.0180064702	27614.8544651066	5.3822306131	1.6642469803	16593.0025965662	0		0
undominated	1	Rx B	36968.4302418399	0	2.95379724	0	0	0		1
	2	Rx B-2	46510.1635413636	9541.7332995238	3.7179836328	0.7641863928	12486.1334747069	0		2
	3	Rx A	74125.0180064702	27614.8544651066	5.3822306131	1.6642469803	16593.0025965662	0		0

CE PSA Output - CE Rankings Report

33.3 Output Distribution graphs

Cost-effectiveness simulations generate many outputs, including cost and effectiveness for each strategy. For each of these outputs, you can generate a probability distribution histogram.

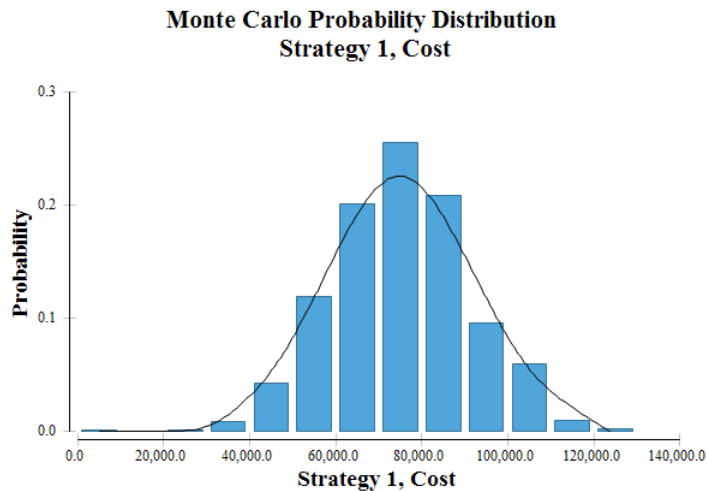
In cost-effectiveness simulations performed at decision nodes, distributions are available not only for each strategy's cost and effectiveness values, but for incremental values as well. Incremental distribution graphs compare one strategy to a baseline strategy.

For cost-effectiveness models, there are multiple views for *Output Distributions* charts.

- *Cost*: Cost Expected Values (EVs) for each strategy.
- *Incremental Cost*: Incremental cost among strategies.
- *Effect*: Effectiveness EVs for each strategy.

- *Incremental Eff*: Incremental effectiveness among strategies.
- *ICER*: Incremental cost-effectiveness ratio among strategies.
- *Net Monetary Benefits*: Net monetary benefits for each strategy for a specified WTP.

When cost or effectiveness is considered in isolation, each view functions like a single EV value for that strategy. Therefore, the *Cost* and *Effect* options generate output similar to the simple tree's EV output. The *Cost* option is shown below.



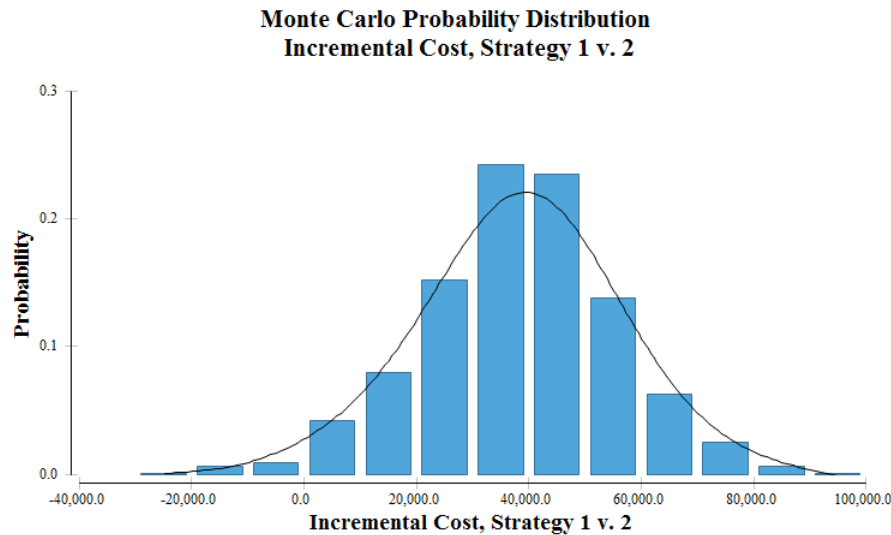
CE PSA Output - Output Distributions - Cost

However, incremental values are quite different. Each incremental histogram allows you to select one strategy as the comparator (first) and another as a baseline (second).

- ▼ Output Distributions ...
 - ▶ ... Cost
 - ▼ ... Incremental Cost
 - [Rx A v. Rx B](#)
 - [Rx A v. Rx B-2](#)
 - [Rx B v. Rx A](#)
 - [Rx B v. Rx B-2](#)
 - [Rx B-2 v. Rx A](#)
 - [Rx B-2 v. Rx B](#)
 - ▶ ... Effect
 - ▶ ... Incremental Eff
 - ▶ ... ICER
 - ▶ ... Net Monetary Benefits (NMB)

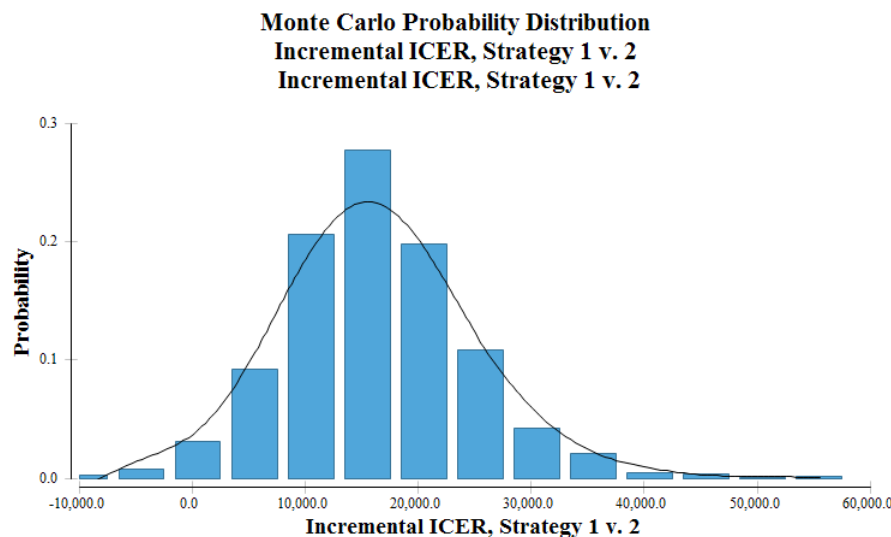
Strategy selections for Incremental Cost graph

For example, selecting the top link compares the Rx A strategy against the baseline of Rx B. Since Rx A is generally a more expensive strategy, the incremental cost will be positive.



CE PSA Output - Output Distributions - Incremental Cost

Although there is natural interest in using the distribution histogram of ICERs to visualize the uncertainty about the cost-effectiveness of strategies, considerable care must be taken in using this graph. The same issues described in the previous chapter, complicating the interpretation of ICER thresholds in sensitivity analysis graphs, also apply in Monte Carlo simulations. It is critical that the scatterplots, acceptability curves, and net benefits graphs and histograms described below are used to put the ICER distribution graph in context — in particular, to determine whether there is uncertainty about the sign (positive or negative) of incremental effectiveness.



CE PSA Output - Output Distributions - ICER

In addition to reading about the other types of CE simulation graphs and reports below, you can read more about the issues surrounding uncertainty analysis and incremental CE ratios in one of the references given at the beginning of the Cost-Effectiveness Modeling and Analysis Chapter.

Histograms of the distribution of net monetary benefit values for a single option in a cost-effectiveness simulation are generated based on a single WTP value. Before generating a histogram, TreeAge Pro will prompt you to enter the WTP.

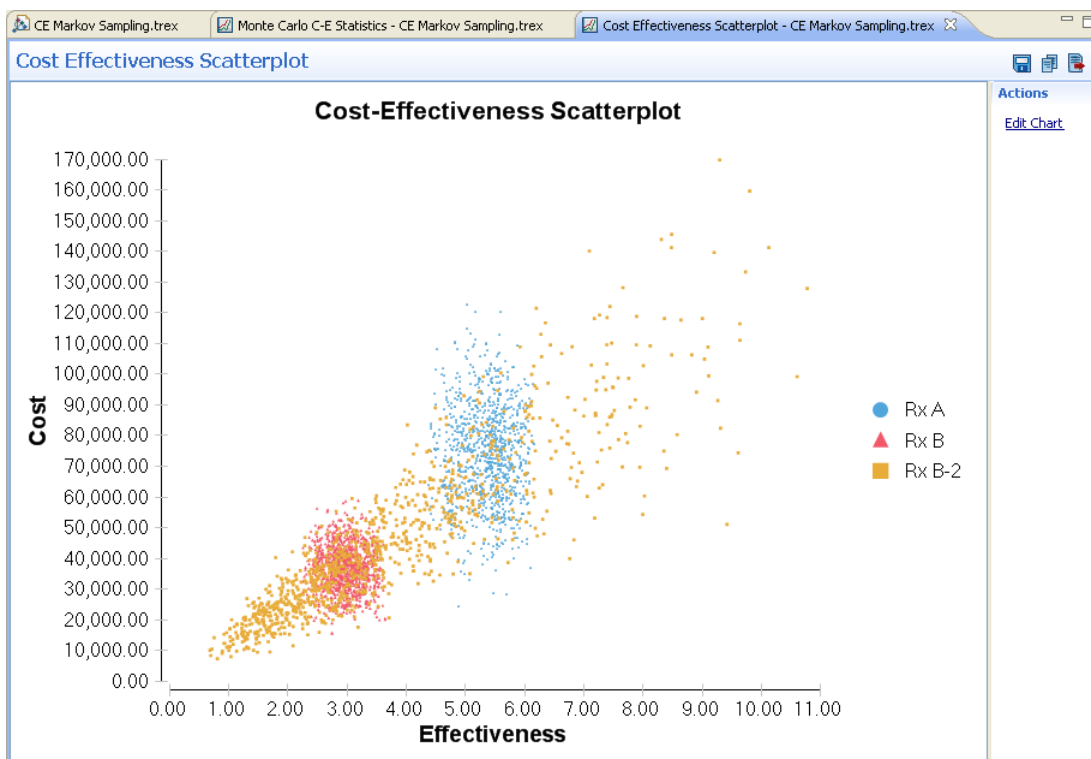
33.4 Scatterplots

The CE simulation output generates two types of scatterplot graphs:

- *CE Scatterplot*
- *ICE Scatter + Ellipses*

33.4.1 CE Scatterplot

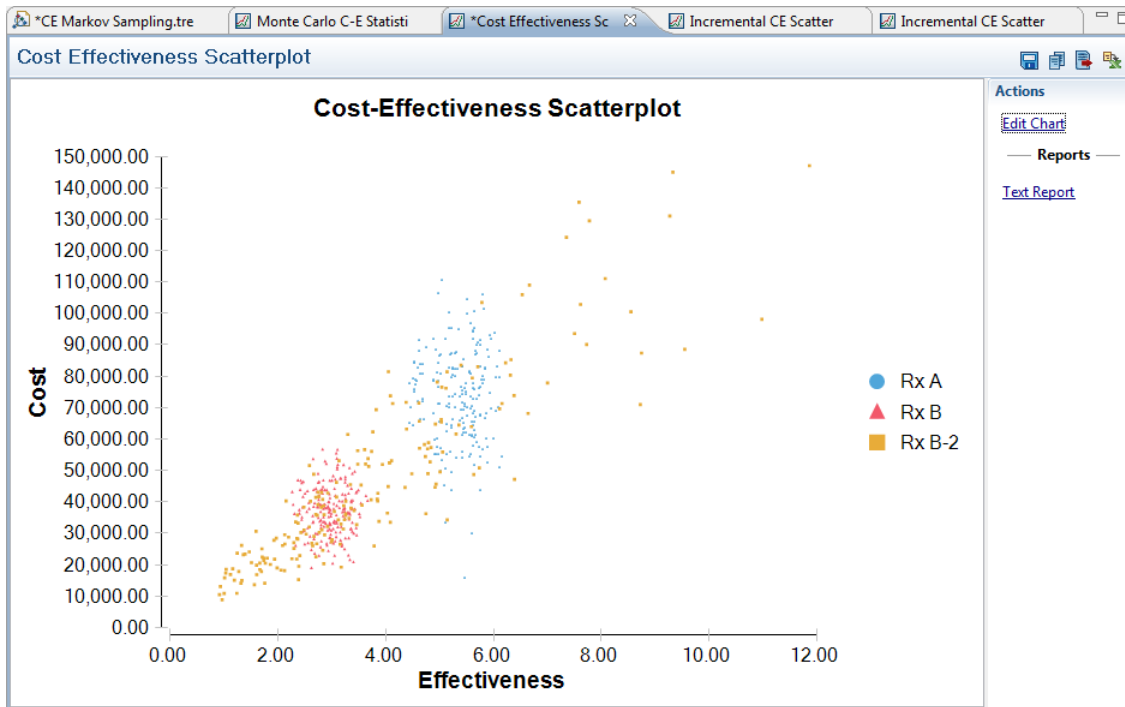
The CE graph, which plots the mean cost and mean effectiveness of each strategy, can be naturally extended to a scatterplot for the simulation. The *CE Scatterplot* uses the cost-effectiveness plane to plot the individual cost and effectiveness pairs for each recalculation of the model. If the simulation is performed at a decision node, each strategy's set of points uses a different color.



CE PSA Output - CE Scatterplot

Depending on the number of iterations included in the simulation, it may be useful to either include only a subset of results in the plot (e.g., if the general density of points is too high) or to increase the size of the dots used in the plot (e.g., if there are only a few points to display). These techniques are described

in the Customizing scatterplot graphs section of the Graph Windows Chapter. The graph below uses the same output, but only 20% of the scatterplot points.



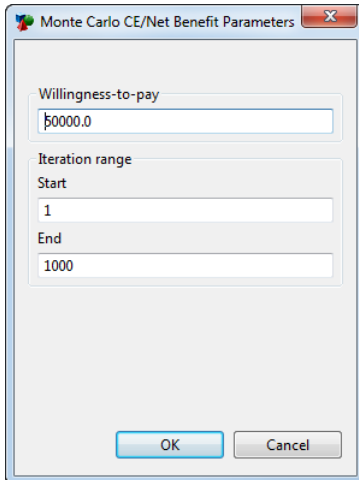
CE PSA Output - CE Scatterplot (filtered)

33.4.2 ICE Scatterplot

Like the CE Scatterplot, the *ICE Scatterplot* uses a form of the standard CE plane to plot points for each iteration in the simulation output. The ICE scatterplot includes a single set of points representing pairs of incremental cost and effectiveness values from the simulation results, based on a comparator (e.g., Option C) relative to a baseline (e.g., Option B). You must select the comparator and baseline strategies via the appropriate report link (comparator first and baseline second).

Generally, if an option has a higher mean cost and effectiveness in the simulation statistical summary, it should be specified as the comparator. The points in the scatterplot will represent the comparator's incremental cost and incremental effectiveness relative to the baseline (represented by the origin). If you select the opposite comparator/baseline combination, most of the incremental values will be negative, making it harder to interpret the graph.

After you click on an ICE scatter link, a dialog box is presented to allow easy access to the WTP and number of iterations for the graph.



Monte Carlo CE/Net Benefit Parameters

Willingness-to-pay
\$00000.0

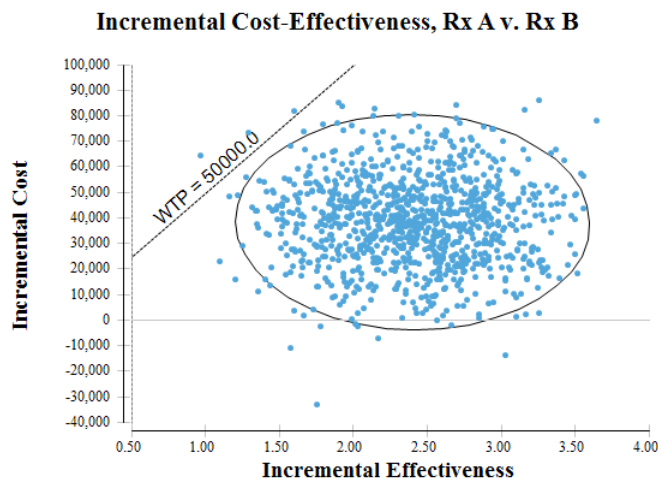
Iteration range
Start
1
End
1000

OK Cancel

CE PSA Output - ICE Scatterplot dialog

The WTP, or ceiling ICER, is used as the slope of a line intersecting the origin of the plot. The WTP line in the graph intersects points having the specified ICER value, and the region below the line includes cost-effective points. This is utilized in the scatterplot's Text Report and ICE Report to calculate the percentage of simulation iterations for which the comparator is cost-effective. A similar analysis is done using a different graph, the Acceptability Curve, which works for all strategies in the analysis, not just two.

The resulting ICE Scatterplot for Rx A v. Rx B is presented below with incremental effectiveness (IE) on the x-axis and incremental cost (IC) on the y-axis. Note the ellipsis showing the 95% confidence interval.



CE PSA Output - ICE Scatterplot graph

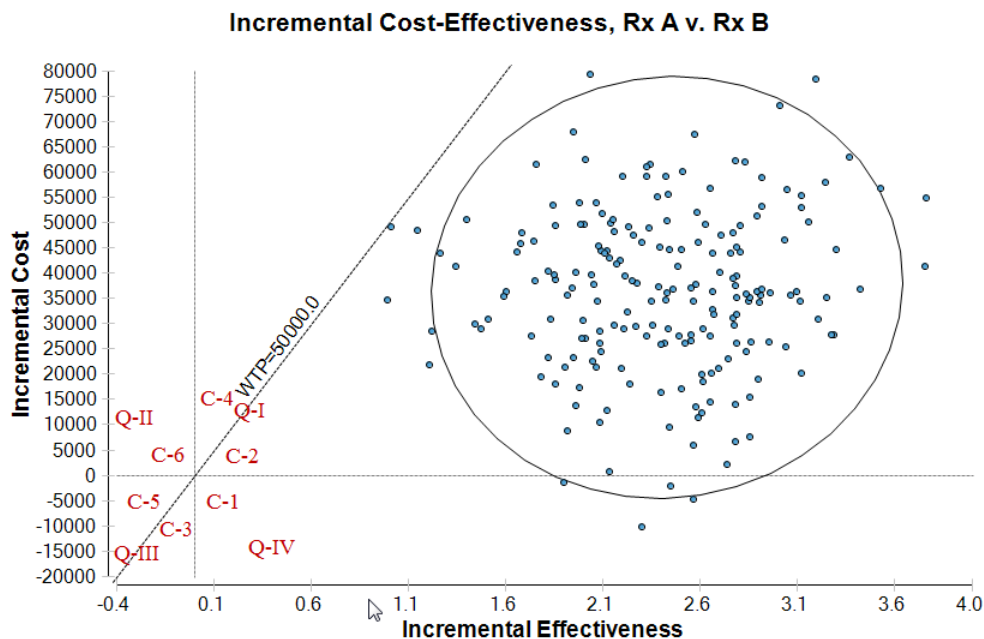


For details on how confidence ellipses account for correlation between cost and effects, see “Reflecting Uncertainty in Cost-Effectiveness Analysis”, Manning et al, Ch. 8 in *Cost-Effectiveness in Health and Medicine*, Gold et al., Oxford Univ. Press (1996).

You can customize the scatterplot after it is generated using techniques described in the Customizing scatterplot graphs section of the Graph Windows Chapter. Customizations include:

- Changing the WTP.
- Changing the confidence interval.
- Changing the scale for the axis.
- Changing the marker type and/or size.
- Filtering the number of scatterplot points.

The graph below uses the same output, modified to show only 20% of the scatterplot points, to set the x-axis scale to include the 0 value and to show smaller markers.



CE PSA Output - ICE Scatterplot graph (customized)

Though not part of the graph itself, the graph's quadrants and components are identified in red. Each component implies a relationship between the two strategies based on IC and IE relative to the ICER threshold. The components are shown in the ICE Report described later in this section.

The graph's Text Report shows the IC and IE for each iteration, which becomes the source of the scatterplot. See below.

Incremental Cost Effectiveness									
ITERATION	COST1	COST2	EFF1	EFF2	COMPARATOR	BASELINE	incrCost	incrEff	
1	61973.2884206899	33984.667961745	5.1082922655	2.9283865737	1	2	27988.620458945	2.1799056918	
2	66078.0199679759	36769.6911316765	5.3476267592	2.5201234509	1	2	29308.3288362994	2.8275033083	
3	78923.6149146909	28289.5792783808	5.8091133998	3.1194036175	1	2	50634.0356363101	2.6897097823	
4	92798.5571472668	31760.1975103173	4.9357496616	2.5243876874	1	2	61038.3596369495	2.4113619741	
5	76526.192624552	29783.8518292768	5.3770263372	2.8652118261	1	2	46742.3407952752	2.5118145112	
6	64119.4313665364	55327.30646636	5.2849230185	2.6208955333	1	2	8792.1249001765	2.6640274853	
7	69399.9546538714	47295.2350724903	5.60137358	2.736300773	1	2	22104.7195813812	2.865072807	

CE PSA Output - ICE Scatterplot Text Report

The graph's ICE Report summarizes the graph's individual data points by component. See below.

Incremental CE Plot Report						
COMPONENT	QUADRANT	INCREFF	INCRCE	INCRCE	FREQUENCY	PROPORTION
C1	IV	IE>0	IC<0	Superior	8	0.008
C2	I	IE>0	IC>0	ICER<50000.0	988	0.988
C3	III	IE<0	IC<0	ICER>50000.0	0	0
C4	I	IE>0	IC>0	ICER>50000.0	4	0.004
C5	III	IE<0	IC<0	ICER<50000.0	0	0
C6	II	IE<0	IC>0	Inferior	0	0
Indiff	origin	IE=0	IC=0	0/0	0	0

CE PSA Output - ICE Scatterplot ICE Report

This summary can be interpreted to indicate the number of iterations that recommend the comparator strategy over the baseline strategy as follows:

- C1 - Comparator is less costly and more effective. Comparator is recommended because it absolutely dominates baseline.
- C2 - Comparator is more costly and more effective. Comparator is recommended because the ICER does not exceed the WTP.
- C3 - Comparator is less costly and less effective. Comparator is recommended because the ICER does not exceed the WTP.
- C4 - Comparator is more costly and more effective. Comparator is not recommended because the ICER exceeds the WTP.
- C5 - Comparator is less costly and less effective. Comparator is not recommended because the ICER exceeds the WTP.
- C6 - Comparator is more costly and less effective. Comparator is not recommended because it is absolutely dominated by the baseline.

Increasing the WTP value changes the slope of the WTP line, the shape of the component regions #2–5, and thus the report. See the section on Acceptability Curves below for a robust method of testing a range of WTP values for all potentially cost-effective strategies.

For additional discussion, refer to "Uncertainty in Decision Models Analyzing Cost-Effectiveness," Maria Hunink et al, *Medical Decision Making* 18:337-346 (1998).

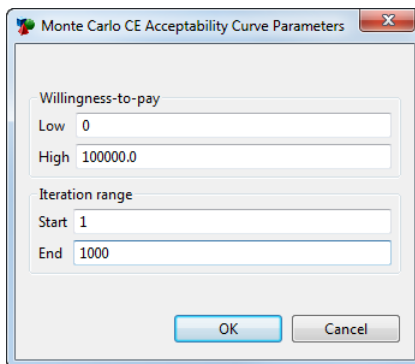
33.5 Acceptability and Net Benefits curves

The *Acceptability Curve* is a commonly-used visual aid for communicating the results of probabilistic sensitivity analysis in cost-effectiveness models.

TreeAge Pro's Acceptability Curve presents the relative cost-effectiveness as a function of the ICER threshold (WTP). For each WTP value, the graph uses net benefits to determine the percentage of simulation iterations that favor each strategy. The percentages will increase for more effective strategies as the WTP increases.

To generate the Acceptability Curve from CE PSA simulation output:

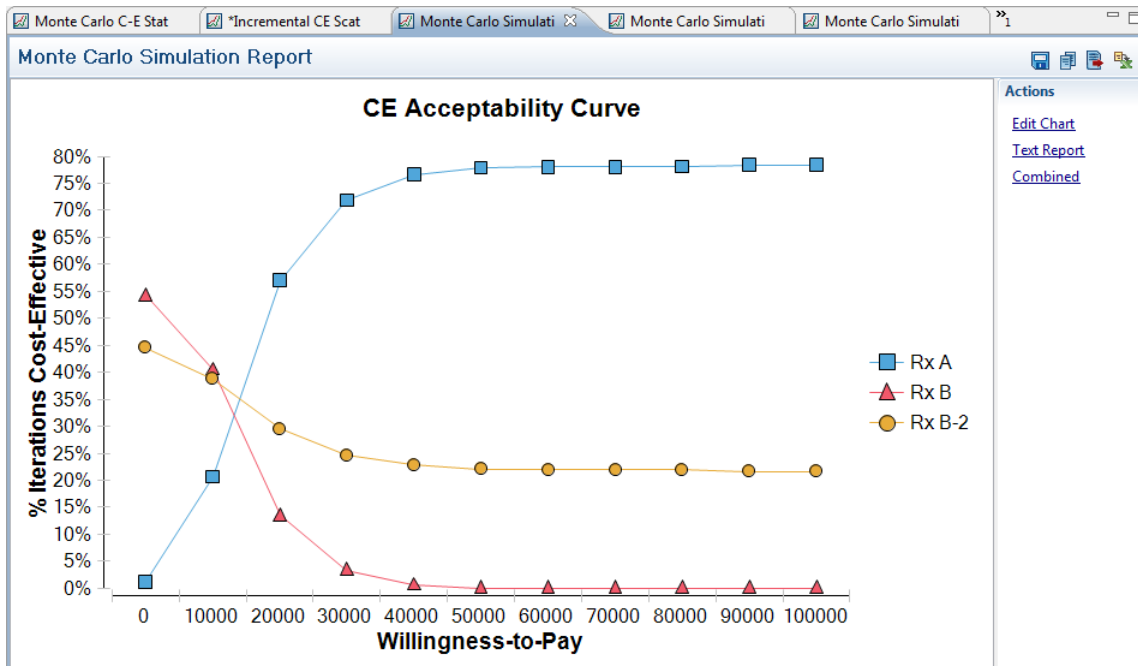
- Select the graphical output Plots/Curves > CE Acceptability > Acceptability Curve.
- Enter the curve's parameters in the Monte Carlo CE Acceptability Curve Parameters dialog (see below). Click OK.



Monte Carlo CE Acceptability Curve Parameters Dialog

Like a sensitivity analysis, the Acceptability Curve requires a range of values for the threshold ICER. You can also select the range of iterations to include in the graph. It is recommended that you select all iterations.

The net benefits version of the Acceptability Curve includes a line for each strategy, with the curves summing to 100% at each interval.



Acceptability Curve

The interpretation usually applied to the net benefits acceptability curves is that the graphed value of any comparator at a particular WTP represents the probability that it is cost-effective (most effective option within the threshold ICER), based on the uncertainties included in the simulation.

The value of a comparator at WTP=0 represents the probability that it is the least costly option. Provided you have entered a sufficiently large top WTP value, the strategy with the highest percentage furthest to the right represents the probability that it is the most effective option.

For more information on the net benefits acceptability curve, refer to "Quantifying stochastic uncertainty" Glick, Briggs, and Polsky, *Expert Rev Pharmacoeconomic Out Res* 1(1), 25-36 (2001) [and www.future-drugs.com].

The Acceptability Curve's Text Report provides the numerical data for each option's probabilities at each WTP interval.

WEIGHT	STRATEGY	STRATEGYNAME	ACCEPTABILITY
0	0	Rx A	0.012
0	1	Rx B	0.542
0	2	Rx B-2	0.446
10000	0	Rx A	0.207
10000	1	Rx B	0.405
10000	2	Rx B-2	0.388
20000	0	Rx A	0.57
20000	1	Rx B	0.135
20000	2	Rx B-2	0.295
30000	0	Rx A	0.719
30000	1	Rx B	0.034
30000	2	Rx B-2	0.247
40000	0	Rx A	0.766
40000	1	Rx B	0.006
40000	2	Rx B-2	0.228
50000	0	Rx A	0.779

Acceptability Curve Text Report

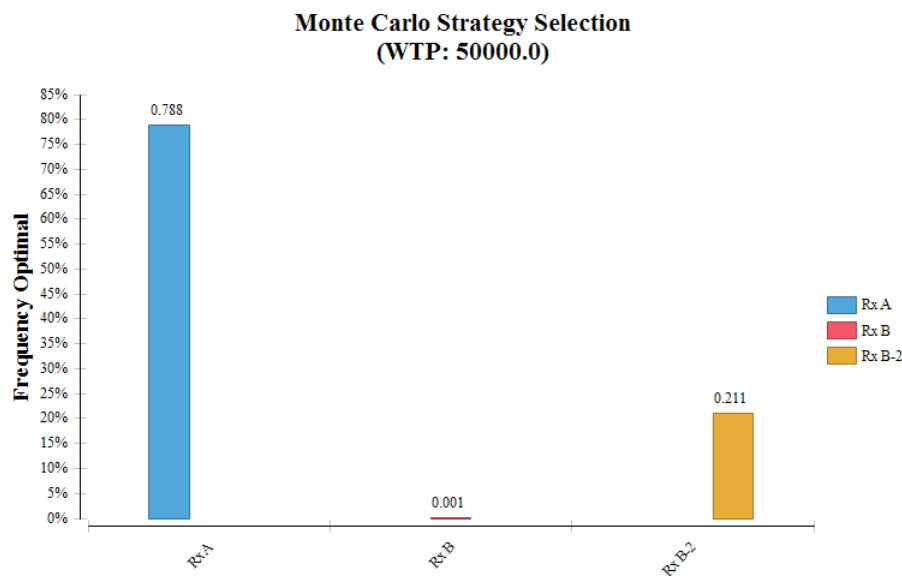
The Combine option to the right of the graph allows you to show both the graph and the text report in the same output window.

33.5.1 Acceptability Frontier

The Acceptability Frontier graph has not yet been implemented in TreeAge Pro 201x.

33.5.2 Strategy Selection / Acceptability Curve

The *Strategy Selection / Acceptability Curve* option shows the percentage of iterations that favor each strategy based on net benefits calculations at a specific willingness to pay value (rather than a range).

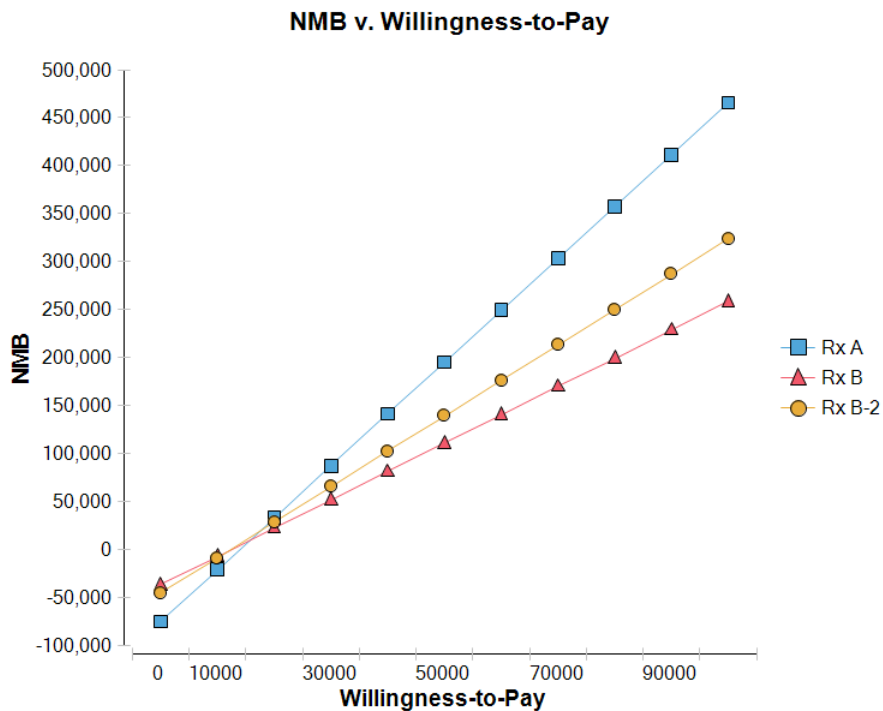


Strategy Selection/Acceptability Curve Graph

33.5.3 Net Benefits vs. WTP

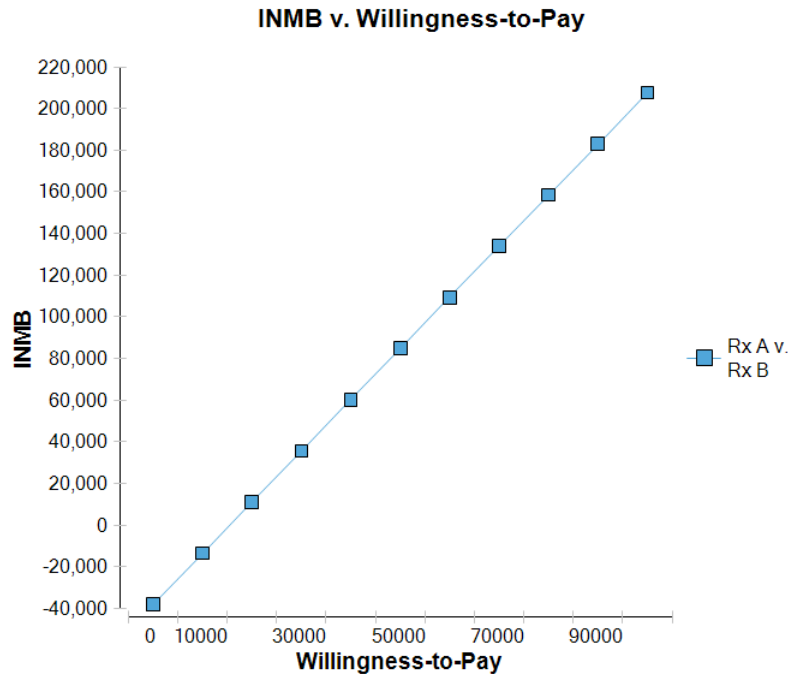
Net monetary benefits (NMB) calculations integrate a particular WTP value and will prompt you to provide a WTP value range. The *Net Benefits vs. WTP* graph, in both average and incremental formats, functions as a sensitivity analysis on WTP.

An intervention's mean effectiveness and cost statistics for the simulation are the only other inputs for each calculation of that intervention's net benefit.



Net Benefits vs. WTP

To graph incremental net benefit curves for any combination of two strategies, click on the appropriate Incremental (INMB) v. WTP link.



Incremental Net Benefits vs. WTP

Incremental net health benefit (INMB) is calculated as:

$$INMB_{C-B} = NMB_C - NMB_B$$

where C refers to a comparator and B refers to the baseline. An alternative form of the equation, providing the same result, is:

$$INMB_{C-B} = ((E_C - E_B) * WTP) - (C_C - C_B)$$

Note that the incremental net benefits graph above shows that INMB is zero at about $WTP = 15,500$. This threshold is also reflected in the Net Benefits vs. WTP graph further above with the crossing of the two strategies' curves.

33.6 Cost-effectiveness value of information (EVPI and EVPPI)

Using the results of a Monte Carlo simulation performed at a decision node, TreeAge Pro can calculate the expected value of perfect information (EVPI) or partial EVPI (EVPPI), based on hypothetically eliminating the simulated uncertainties (normally, one or more sampling distributions).

To generate an EVPI/EVPPI summary report:

- Run a CE PSA simulation at a decision node (as was done earlier in this chapter).
- Expand the Text Reports group to the right of the simulation summary data.
- Click on the EVPI\EVPPI Summary Report link.
- Enter the WTP value and the iteration range.

You will be presented with the following EVPI/EVPPI summary report.

Weight (WTP)	EVPI (INMB)	Avg. IC with PI	Avg. IE with PI	Optimal Strategy
40000	9,790.51	-1,201.49	0.215	1

EVPI/EVPPI Summary Report

The summary report reflects the overall mean values generated by the EVPI/EVPPI value calculated for each iteration in the simulation. Let's look at the EVPI/EVPPI Details Report (via link shown above) to see how the values are calculated for each iteration.

Iteration	Weight (WTP)	EVPI (Incr. NMB)	Incr. Cost with PI	Incr. Eff with PI	Iteration Best Strategy	Cost (Iteration Strategy)	Eff. (Iteration Strategy)	Simulation Optimal Strategy	Cost (Optimal Strategy)	Eff. (Optimal Strategy)
1	40000	0	0	0	1	60845.9105765672	5.1994994022	1	60845.9105765672	5.1994994022
2	40000	0	0	0	1	49346.4598929069	5.092094772	1	49346.4598929069	5.092094772
3	40000	0	0	0	1	69820.790242679	5.5795581006	1	69820.790242679	5.5795581006
4	40000	5980.0101188886	-8570.4668090083	-0.0647614173	3	83723.6799409025	5.6844395666	1	92294.1467499108	5.74920098
5	40000	12448.0201049...	-1911.0137526674	0.2634251588	3	77570.0626926064	6.0316299762	1	79481.0764452738	5.76820481
6	40000	0	0	0	1	86784.6164993528	5.743176027	1	86784.6164993528	5.743176027
7	40000	0	0	0	1	92882.0876093594	4.963931119	1	92882.0876093594	4.963931119
8	40000	0	0	0	1	79827.1441253895	5.7382096767	1	79827.1441253895	5.7382096767
9	40000	0	0	0	1	72461.7288049042	5.1042273979	1	72461.7288049042	5.1042273979
10	40000	0	0	0	1	46721.3339954708	5.19041921	1	46721.3339954708	5.19041921
11	40000	0	0	0	1	57772.330296551	5.8012098127	1	57772.330296551	5.8012098127
12	40000	43453.954194247	-42876.9735042...	0.0144245173	3	53648.1914997588	4.9773942903	1	96525.1650040038	4.96296977
13	40000	0	0	0	1	63368.8030973644	4.8286139083	1	63368.8030973644	4.8286139083
14	40000	0	0	0	1	56122.0934894579	5.8421313949	1	56122.0934894579	5.8421313949
15	40000	0	0	0	1	58410.2197702693	5.5386563224	1	58410.2197702693	5.5386563224
16	40000	0	0	0	1	73363.8680835652	4.8843069276	1	73363.8680835652	4.8843069276
17	40000	0	0	0	1	73390.600085446	5.3953336038	1	73390.600085446	5.3953336038
18	40000	0	0	0	1	63390.6519138907	5.8401351523	1	63390.6519138907	5.8401351523
19	40000	27635.5317262...	-60409.2646445...	-0.819343323	3	39126.8446265033	4.1364664716	1	99536.1092710585	4.95580975

EVPI/EVPPI Details Report

The EVPI/EVPPI reports use net monetary benefit calculations (NMB) using the WTP parameter specified when running the report. Each strategy's mean cost and effectiveness values are used to determine the overall optimal strategy for that WTP. Then, separate calculations are done to determine the optimal strategy for each of the simulation's iterations.

When the iteration's optimal strategy is the same as the overall optimal strategy, their EVPI/EVPPI value is zero. However, when the iteration's optimal strategy is different, then there is some value to eliminating the uncertainties being simulated. The EVPI (incremental NMB) is equal to the NMB value for the iteration's optimal strategy, less the NMB value for the overall optimal strategy.

The average, or expected value of information for all iterations, is reported in the summary report. In the example above, the expected cost savings from perfect information (per patient) is 1,201.49 and the expected gain in effectiveness is 0.215.



The description of these reports refers to EVPI and EVPPI.

EVPI is calculated when running regular probabilistic sensitivity analysis with a single parameter sampling loop (and possibly also a microsimulation/trials loop).

EVPPI is calculated when running two probabilistic sensitivity analysis parameter sampling loops. This allows you to isolate one or more distributions in the outer-most loop to isolate the effect of EVPPI for those specific parameter uncertainties.

34. Building and Analyzing Markov Models

This chapter covers the basics of creating and analyzing Markov processes with TreeAge Pro Healthcare and TreeAge Pro Suite. Some basic conceptual background is provided.

The Markov Modeling Tools and Techniques Chapter covers a variety of important Markov topics including: time-dependent probabilities, discounting, half-cycle correction, tunnel states, and Markov decision processes. Markov microsimulation is covered in detail in the Individual-Level Simulation and Markov Models Chapter.

34.1 Markov modeling basics

While most decision trees include a simple notion of time (i.e., left to right chronologically), there are no shortcuts in a standard tree structure for representing events that recur over time. A state transition model, also called a Markov model, is designed to do just this. Markov models are used to simulate both short-term processes (e.g., development of a tumor) and long-term processes (e.g., an individual's lifespan).

34.1.1 State transition models

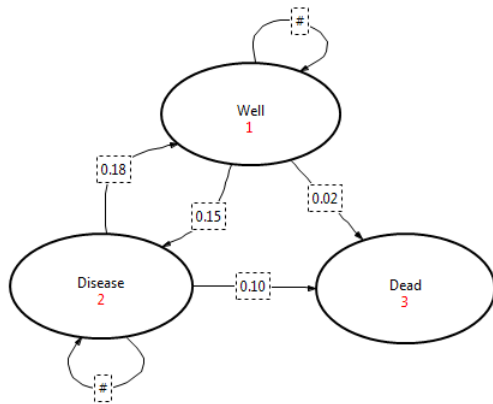
Markov models built in TreeAge Pro often represent discrete-time state transition models (although discrete event modeling is also possible). A discrete-time Markov model usually follows a basic design, such that:

- The time period of interest (i.e., 10 years) is divided into equal intervals, or *cycles*.
- A finite set of mutually exclusive *states* is defined such that, in any given cycle, a member of the cohort is in only one state.
- *Initial probabilities* determine the distribution of cohort members among the possible states at the start of the process (often, the entire cohort starts in the same state).
- A matrix of *transition probabilities*, applied in each successive cycle, defines the possible changes in state.
- To calculate an expected value for the model, (e.g., net cost or quality-adjusted life expectancy), different cost and/or utility *rewards/tolls* are accumulated for each interval spent in a particular state.

Characteristics of a discrete-time Markov model

34.1.2 Graphical representation

In a state transition diagram (see below), each state is represented using an oval, arrows represent transitions, and numbers along the arrows indicate the transition probabilities. The probabilities of the transition arrows emanating from any state must sum to 1.0.



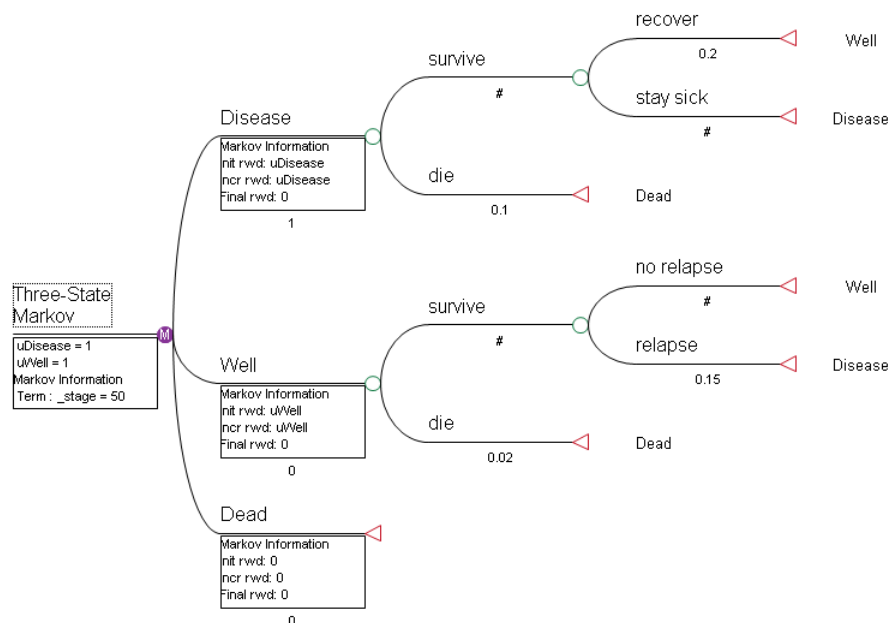
Markov as State Transition Diagram



Click here for information on creating state transition diagrams.

TreeAge Pro represents Markov models in a decision tree format, which is more easily integrated into decision trees.

The Health Care tutorial example tree "Three-State Markov" is presented below.



Three-State Markov Model



Markov cycle trees can be appended to paths in a TreeAge Pro decision tree anywhere you might place a terminal node.

34.1.3 Calculation basics

There are two commonly-used methods for evaluating a Markov model: expected value calculation (called "cohort" analysis), and Monte Carlo simulation (first-order trials or microsimulation). It is

important to understand the difference between the two analysis methods, and to recognize the terms associated with them.

In an *expected value* analysis, the *percentage of a hypothetical cohort* in a state during a cycle is multiplied by the cost or utility associated with that state. These products are summed over all states and all cycles. In TreeAge Pro, expected value calculations are the basis of most analyses, including *n*-way sensitivity analysis and baseline cost-effectiveness analysis.

On the other hand, in a *microsimulation* (a.k.a., discrete simulation), a single trial's value is simply the sum of the rewards/tolls/payoffs for the path traversed by an "individual" taking a random walk through the model's chance nodes (using a Monte Carlo pseudo-random number series). An expected value is estimated by averaging as many trials as possible.

In TreeAge Pro, the same Markov model can be evaluated by either expected value or simulation methods. Generally deterministic, expected value analysis is preferred because it is more computationally efficient; it returns a mean value much more quickly than simulation, which often requires thousands of trials to return a mean value within an acceptable error. However, some models will require simulation; refer to the Individual-Level Simulation and Markov Models Chapter.

Additional background discussion can be found in:

- *Decision Making in Health and Medicine*, Hunink, and Glasziou (2001), Cambridge University.

You are urged to consult a variety of publications dealing with the concepts which underlie Markov modeling and simulation.

34.1.4 Non-standard Markov models

In TreeAge Pro, the basic Markov modeling rules outlined above can be overruled in a variety of ways, for example:

- Time-dependent Markov models are easily handled using tables, tunnels, and/or tracker variables.
- Discrete event models can combine samplings from event time distributions and simulation features like tracker variables, parallel trials, and dynamic populations.
- A Markov model can be analyzed using the Node() function in such a way that sensitivity analysis and other cohort-type analyses can be used, while the Markov model is actually evaluated using microsimulation trials.
- EV/cohort analysis of a Markov model can make use of a dynamic cohort with a specific starting size and composition that may change over time.

Exceptions to Markov modeling rules supported by TreeAge Pro

These kinds of features are covered in the next few chapters.

34.2 Building a Markov cycle tree in TreeAge Pro

The design of a basic Markov model requires consideration of a number of components, most of which have been introduced above:

- *States* – The set of distinct health states under consideration in the model, together with the possible transitions between them.
- *Cycle length* – The length of time represented by a single stage (or cycle) in a Markov process. This value is implicit in the probabilities, rewards/tolls, and termination condition.
- *Initial probabilities* – A set of probabilities used only at the outset of the process, describing the initial distribution of the cohort among the states.
- *Transition probabilities* – The matrix of probabilities of moving between health states from one stage to the next.
- *Rewards/tolls* – Per-cycle costs and/or utilities (corresponding to payoffs in regular trees) representing the outcome measure(s) being calculated, e.g., costs or QALYs. Rewards/tolls may be associated with health states, or with instantaneous, short-term events.
- *Termination condition* – A logical test evaluated at the beginning of each new cycle to determine if the process should continue or stop.

Components of Markov models in TreeAge Pro

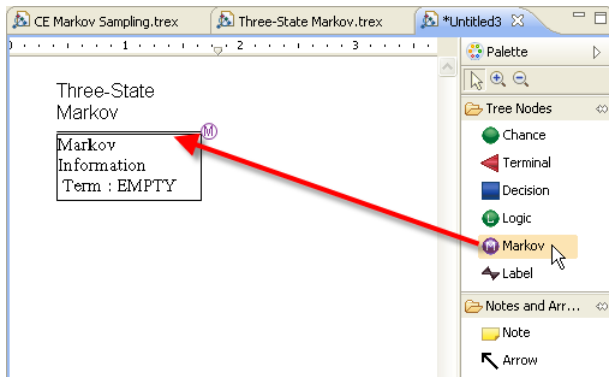
These elements will be illustrated using the simple, three-state Markov state transition model illustrated at the beginning of the chapter.

34.2.1 The Markov node

To begin, a Markov node must be used. Any number of Markov nodes can be included in a decision tree. In this case, the root node of an empty tree will be changed to a Markov node.

To create a Markov node:

- Create a new tree by choosing File > New from the menu.
- Choose Blank Tree Diagram from the template dialog.
- Right-click on the root Decision Node and choose Change Type > Markov from the context menu.
- Type "Three-State Markov" for a text description of the new Markov node.



Create a Markov node

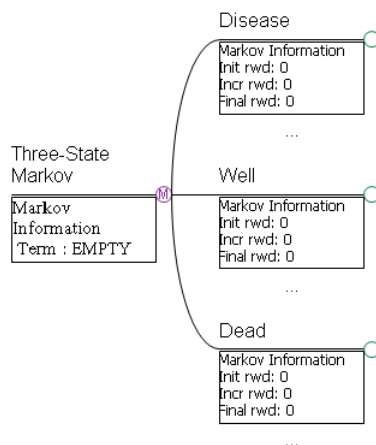
Note that, in addition to using the purple Markov node symbol, TreeAge Pro also adds a Markov information box below the node; we will return to this later.

34.2.2 Markov states (and initial probabilities)

The branches of the Markov node enumerate the Markov states, and are labeled as such. For our simple three-state model, we need three branches from the Markov node.

To add the Markov states:

- Drag three chance nodes from the palette to the Tree Diagram Editor as branches of the Markov node.
- Name the three new nodes *Disease*, *Well* and *Dead* from top to bottom.



Add Markov states

TreeAge Pro's use of arcs, rather than straight lines, for the branches within a Markov subtree is simply to make it easier to distinguish Markov subtrees from the rest of a decision tree.

Below the Markov state branches, *initial state probabilities* must be entered. These probabilities, like the probabilities of a chance node, must sum to one. It is often the case that many states will have initial probabilities of 0. In the three-state model, for example, 100% of the cohort begins in the Disease state.

To enter the initial probabilities for the Markov states:

- Below the branches of the *Disease*, *Well* and *Dead* states, enter the initial probabilities, 1, 0 and 0, respectively.

Initial state probabilities are used by TreeAge Pro only once during the Markov process, to determine where individuals should spend the first cycle of the process. All subsequent movement through the model utilizes *transition probabilities*, which you will specify later.

34.2.3 State rewards

In TreeAge Pro, costs or utilities assigned in a Markov model are called *rewards*. A *state reward* refers to a value that is assigned to individuals because they spend one cycle in a particular state. This might be a cost or a unit of life expectancy, for example.

The state rewards must reflect the *length of a cycle*. For instance, if you have decided on a yearly cost of \$6000 for a particular state, but your cycle length is 1 month (not 1 year), then the state reward (cost) should be $6000/12$, or 500 per cycle.

Assume that the goal is to estimate average life expectancy, and that the model's cycle length is one year. To calculate life expectancy in terms of years, you would assign a state reward of 1 to any alive state. (If the cycle length were 1 month, a state reward equal to $1/12$ would be used to calculate life expectancy in years.)

For life expectancy (or other) calculations, you can enter three separate state reward expressions at each Markov state. The reasons for having three separate state rewards — primarily for the half-cycle correction — will be explained in detail in the next chapter.

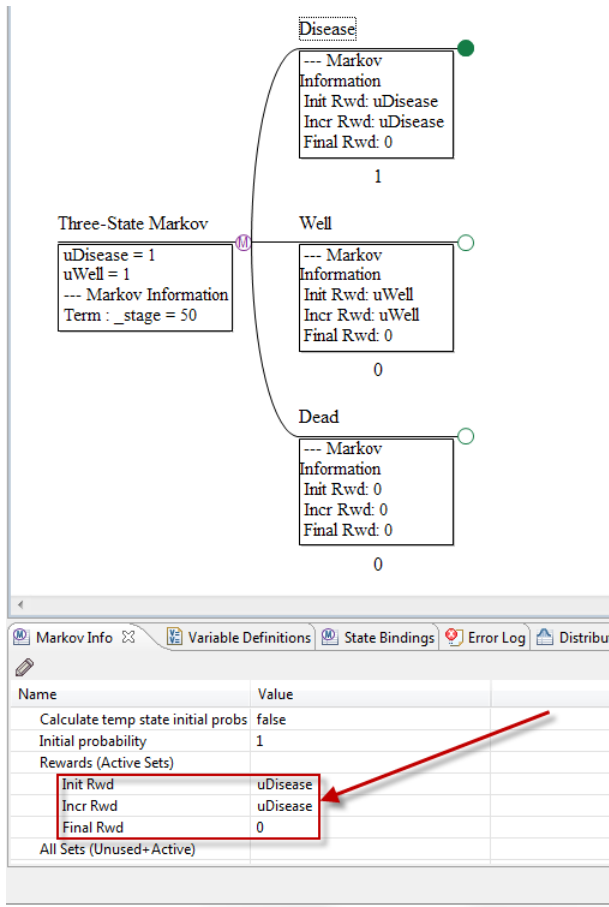
A state's initial reward is assigned only in the first cycle, stage 0, and only to individuals that spend stage 0 in that state. The incremental reward is assigned in subsequent cycles during the process. The final reward (if any) is assigned after the process is over to individuals ending up in that state.

In this case, we want to assign a reward of 1 for each year/cycle someone spends in an alive state. Instead of assigning numeric initial (stage 0) and incremental state rewards of 1 for the Disease and Well states, use a variable to represent the "utility" of each state.

To define state rewards:

- Define the variable *uDisease* at the root node with a value of 1.
- Select the *Disease* node.
- Choose Views > Markov Info from the menu.

- In the Markov Info view enter the Init rwd, Incr rwd and Final rwd values $uDisease$, $uDisease$ and 0, respectively.
- Now enter the state rewards for the other alive Markov state, *Well*, using the values $uWell$, $uWell$ and 0, respectively.
- The zero rewards can be left unchanged in the Dead state.
- Save the partially complete tree as Three-State Markov.



Enter Markov state rewards

The state rewards are also visible within a box under the Markov state node. This display can be turned on and off via the "Show Markov information" option in the Tree Preferences category Variables/Markov Info.

Technically, assigning an initial reward in Well is unnecessary, because its initial probability is 0. However, it does not hurt to specify it as this gives you the flexibility of later changing the initial probability of Well, for instance, in a sensitivity analysis. Similarly, using variables for the rewards provides flexibility for later modifications.

You can also enter Markov rewards via the State Reward Dialog.

To open the State Reward Dialog:

- Select a reward in the Markov View.

- Click on the "pencil" icon in the view's toolbar.

Define Initial Reward, Set #1

Node: Disease Half-Cycle Correct

Initial Reward, Set #1 =	Incremental Reward, Set #1 =	Final Reward, Set #1 =
uDisease	uDisease	0
Insert Expressions: Group: Variables, Functions, Operators, Keywords, [empty lists], Trackers, Tables, Distributions Element:	Insert Expressions: Group: Variables, Functions, Operators, Keywords, [empty lists], Trackers, Tables, Distributions Element:	Insert Expressions: Group: Variables, Functions, Operators, Keywords, [empty lists], Trackers, Tables, Distributions Element:
Calculated value: 1.0	Calculated value: 1.0	Calculated value: 0.0
Definition info:	Definition info:	Definition info:

Markov State Info
Description: Disease
Comment:

OK Cancel

State Reward Dialog

The State Reward Dialog allows you to enter the three state rewards for that state via internal formula editors. It also provides an option to perform half-cycle correction at the top-right corner of the dialog.

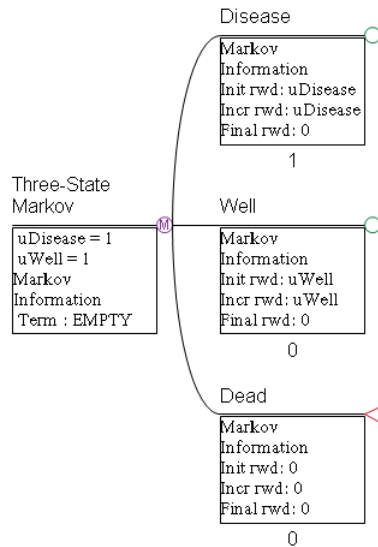
34.2.4 Transition subtrees and absorbing states

The branches of the Markov node represent the possible states. On the other hand, the branches (or subtree) emanating from a Markov state represent possible events during a cycle in that state, including transitions to other states.

The easiest state to complete is *Dead*; a state from which an individual cannot exit has no transition subtree, and is called an absorbing state. To represent an absorbing state, a Markov state is simply changed to a terminal node. Markov models are not required to have any absorbing states. In this example, as in often the case, the Dead state is the sole absorbing state.

To create an absorbing state:

- Right-click on the Dead node and choose Change Type > Terminal from the context menu.



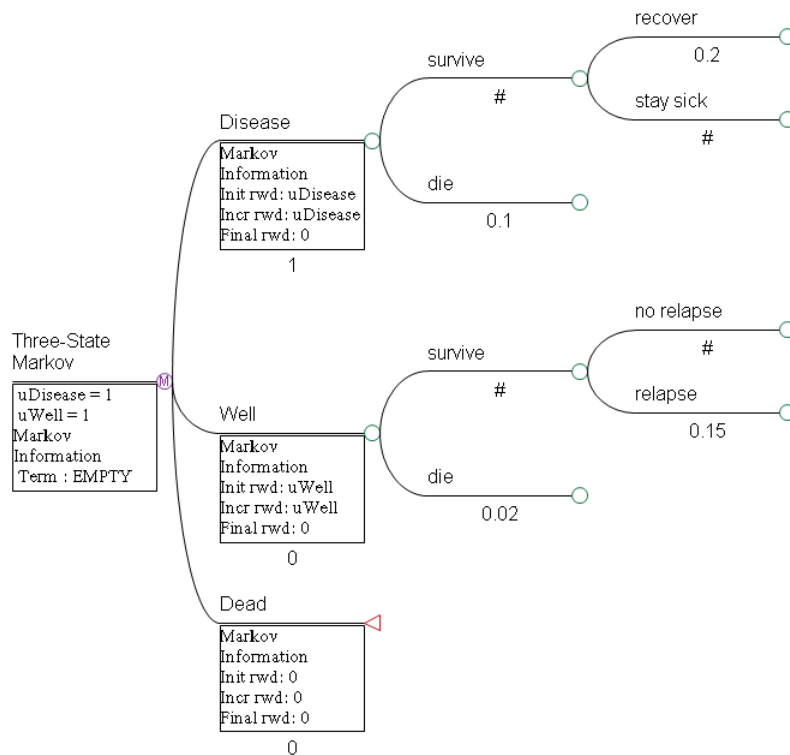
Create absorbing state

Unlike a Markov state transition diagram, the Markov cycle trees that you create in TreeAge Pro can represent a series of events that can occur during a single cycle. Any number of chance nodes, as well as logic and label nodes, can be used to the right of a Markov state.

To illustrate this in the example, the transition subtrees for the *Disease* and *Well* states each use two chance nodes (instead of one chance node with three branches). One chance node represents mortality (including excess mortality in the *Disease* state), and a second represents whether or not an individual gets/stays sick.

To create a transition subtree:

- Right-click on the *Disease* node and choose Add Branch from the context menu. Two chance nodes will be added.
- Label the two new nodes *survive* and *die*.
- Enter the transition probabilities # and 0.1 under the nodes *survive* and *die*, respectively.
- Right-click on the new *survive* node and choose Add Branch from the context menu.
- Label the two new nodes *recover* and *stay sick*.
- Enter the transition probabilities 0.2 and # under the nodes *recover* and *stay sick*, respectively.
- Right-click on the *Well* node and choose Add Branch from the context menu.
- Label the two new nodes *survive* and *die*.
- Enter the transition probabilities # and 0.02 under the nodes *survive* and *die*, respectively.
- Right-click on the new *survive* node and choose Add Branch from the context menu.
- Label the two new nodes *no relapse* and *relapse*.
- Enter the transition probabilities # and 0.15 under the nodes *no relapse* and *relapse*, respectively.



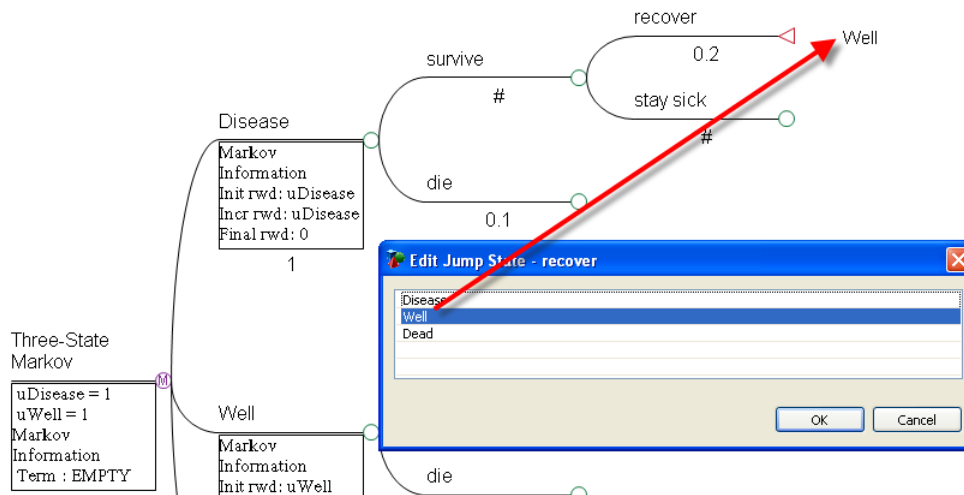
Create a transition subtree

All that is left to do is to terminate each path in the transition subtrees. These transition nodes represent the last event in each path during a cycle, not the end of the Markov process. Individuals reaching a transition node are pointed to a Markov state where they will begin the next cycle (if the process is not terminated first).

To create jump nodes:

- Right-click on the *recover* node and choose Change Type > Terminal from the context menu.
- TreeAge Pro will automatically open the Edit Jump State dialog. In the dialog, select Well from the list of existing states as the appropriate jump-to state for the recover node.
- Click OK to save the jump state and close the dialog.

To the right of each transition node's symbol TreeAge Pro displays the name of the jump-to state for the next cycle.



Add jump states to Markov model

If the wrong jump-to state has been assigned to a transition node, it is easy to change the specified transition.

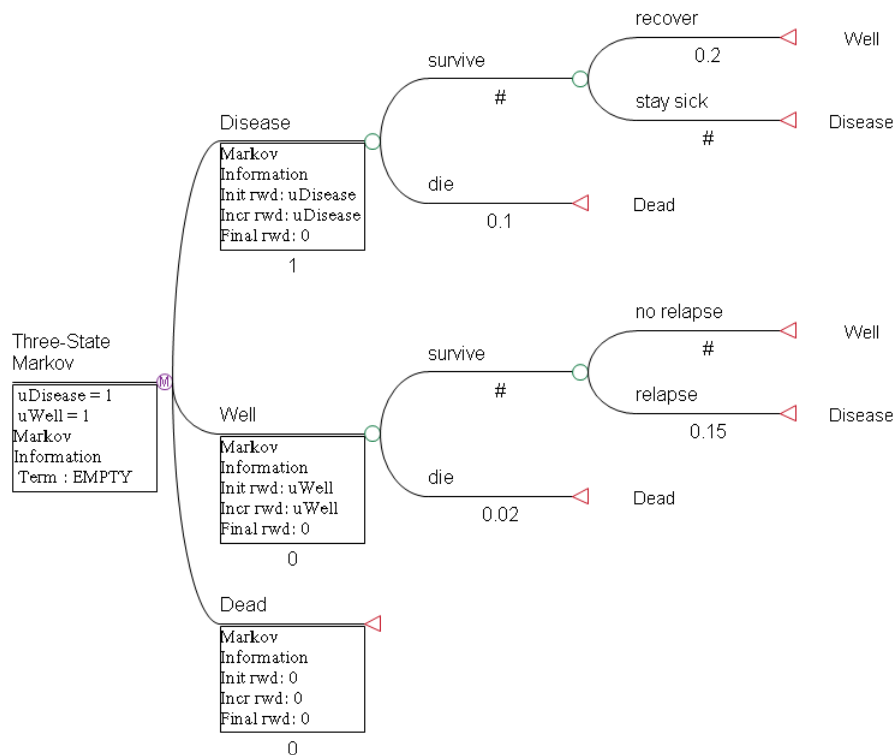
To change a transition node's jump-to state:

- Select the terminal node in the Markov transition subtree.
- Double-click on the jump state to reopen the Edit Jump State Dialog.
- Choose the correct jump state from the list.
- You can also change the jump state in the Markov Info View.



If you change the name of a Markov state, TreeAge Pro automatically updates the transition nodes pointing to it.

Now, set up the remaining jump nodes as seen below. Then save the model.



Markov model with all jump states set

34.2.5 The `_stage` counter and the termination condition

When analyzing a Markov model, TreeAge Pro uses the termination condition, or stopping rule; you specify at the Markov node to determine whether a cohort analysis is complete. TreeAge Pro evaluates the termination condition at the beginning of each cycle except the first. If the condition is *true*, the Markov process ends, final rewards are assigned if necessary, and the results are reported.

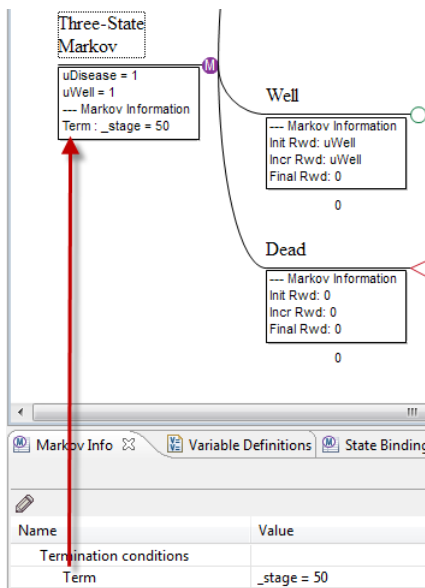
The termination condition is usually very simple. Often, it just checks how many cycles have been completed and stops when a certain number is reached. The number of cycles that have passed is contained in a built-in counter called `_stage`, which TreeAge Pro sets equal to 0 before the first cycle and increments by 1 before each subsequent cycle (i.e., before assigning state rewards).

The termination condition can be a more complex expression, referencing variables, distributions, other keywords, etc. For example, you might want to run until a *variable age* is over 75. In such a case, the following termination condition would be used: `age > 75`. The termination condition can also include multiple conditions, using the logical operators `&`, `|` and `!`, which represent AND, OR and NOT operators respectively. For example, if you wanted the model to run for 50 years or until the age is over 75, the following termination condition would be used: `_stage = 50 | age > 75`

For the Three-State Markov model, use a simple termination condition.

To set the termination condition:

- Select the Markov node.
- Choose Views > Markov Info from the toolbar.
- In the Termination conditions > Term field within the Markov Info View, enter the termination condition `_stage = 50`.
- Save the tree.



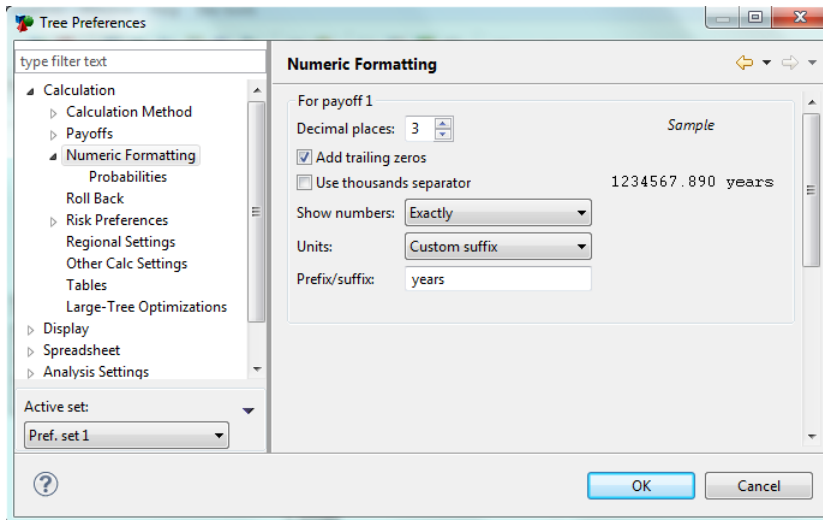
Enter termination condition

The termination condition `_stage = 50` will cause the process to perform 50 cycles (#0 to #49), with an equal number of reward assignments and transitions.

See the next chapter for details on building and interpreting more complex termination conditions.

34.3 Analyzing a Markov model

Once you have completed the Markov model, various kinds of analyses can be performed. Before performing an analysis, however, make sure the tree is set to use appropriate numeric formatting for displaying calculation results. Specifically, change the numeric formatting preferences to match those shown below.



Set numeric formatting preferences

34.3.1 Cohort (expected value) analysis

First, roll back the tree to verify that it is ready to calculate.

To roll back the tree:

- Choose Analysis > Roll Back from the menu.

If you have forgotten to perform one of the steps in the Markov modeling tutorial, an error may be reported identifying the problem node. If there are no errors, TreeAge Pro will display the results of the cohort, expected value calculations on the face of the tree.

Next to the Markov node a roll back box should display an expected value of about 15.891 years (within the 50 years the model is allowed to run). How this value is calculated will be explained below. The boxes next to the Markov states display the average time spent in the state, along with the final probability (FP) of that state when the process terminates. These numbers correspond to the last row of the cohort analysis text report, explained below.

If roll back is still on, turn it off before continuing with another analysis.

To turn off roll back:

- Choose Analysis > Roll Back from the menu.

Usually the preferred expected value analysis for a Markov model is a Markov cohort analysis at the Markov node. It provides detailed text report options, and a variety of graphical outputs.

To perform a cohort expected value analysis (quick):

- Select the Markov node.
- Choose Analysis > Markov Cohort > Markov Cohort (Quick).

After the cohort analysis terminates, the Markov Cohort Analysis output is displayed.

The screenshot shows a software window titled "Markov Cohort (Quick)". It displays a table with the following data:

Stage	State	Probability	State Reward	Stage Reward	Cumulative Reward
Stage 0	Disease	1	1	1	1
	Well	0	0		
	Dead	0	0		
Stage 1	Disease	0.72	0.72	0.9	1.9
	Well	0.18	0.18		
	Dead	0.1	0		
Stage 2	Disease	0.54486	0.54486	0.8244	2.7244
	Well	0.27954	0.27954		
	Dead	0.1756	0		
Stage 3	Disease	0.43339	0.43339	0.76432	3.48872
	Well	0.33093	0.33093		
	Dead	0.23568	0		

On the right side of the window, there is an "Actions" panel with links for "Text Reports" (Summary Report) and "Charts" (State Prob., Survival Curve, State Reward, Stage Reward, Cumulative Reward).

Markov Cohort Analysis Output (Quick)

The output shows the accumulation of rewards by cycle, stage and state. The stage-level columns are:

- *Stage*: the *_stage* (cycle) counter starting with zero.
- *Stage reward*: the reward amount accumulated within that stage.
- *Cumulative reward*: the reward amount accumulated from the first stage through this stage.

The output also includes collapsible stage groupings that show state-level data within the stage. The stage-level columns are:

- *State*: The Markov state.
- *Probability*: The percentage of the cohort starting the stage in that state.
- *State reward*: The reward accumulated in that stage and that state.

The sum of the state rewards will be equal to the total stage reward for the cycle.

Markov Cohort Analysis accumulates state rewards by stage and by state as follows...

- For each stage/state, the *state reward* is the product of the *state probability* (the portion of the cohort starting the cycle in that state) and the *state reward value from the tree* (initial for the first cycle, incremental for subsequent cycles).
- The *stage reward* is the sum of the state rewards for each state.
- The overall *expected value* for the Markov model is the sum of all the stage rewards.

State reward calculation

Let's look at a few of the calculations.

At the beginning of Stage 0, 100% of the cohort is in the *Disease* state, so the state probability is 1. The state reward entered for the *Disease* state is *uDisease*, which is defined as 1. Therefore the Probability value for Stage 0, State *Disease* is 1, as seen below.

$$\langle \text{stage state rwd} \rangle = \langle \text{state prob} \rangle * \langle \text{state rwd from tree} \rangle = 1 * 1 = 1$$

The overall stage reward is equal to that stage's reward for the Disease state since the entire cohort started the cycle in that state.

At the beginning of Stage 1, we start to see the cohort split among all the states.

For Stage 1, state *Disease*:

$$\langle \text{stage state rwd} \rangle = 0.72 * 1 = 0.72$$

For Stage 1, state *Well*:

$$\langle \text{stage state rwd} \rangle = 0.18 * 1 = 0.18$$

For Stage 1, state *Dead*:

$$\langle \text{stage state rwd} \rangle = 0.1 * 0 = 0$$

The total stage reward is the sum of the state rewards for the three states.

$$\langle \text{stage rwd} \rangle = 0.72 + 0.18 + 0 = 0.9$$

When collapsed, the Markov Cohort output shows only the accumulation of rewards by stage. Note that there is a line for Stage 50, even though the analysis is terminated before starting that state. The rewards for that stage would be non-zero if we had entered final rewards for any of the states.

Stage	State	Probability	State Reward	Stage Reward	Cumulative Reward
Stage 35				0.1359170902	14.580145727
Stage 36				0.1289443096	14.7090900366
Stage 37				0.1223292449	14.8314192815
Stage 38				0.1160535447	14.9474728262
Stage 39				0.1100997988	15.057572625
Stage 40				0.1044514904	15.1620241154
Stage 41				0.0990929499	15.2611170653
Stage 42				0.0940093117	15.355126377
Stage 43				0.089186473	15.44431285
Stage 44				0.0846110541	15.5289239041
Stage 45				0.0802703621	15.6091942661
Stage 46				0.076152355	15.6853466211
Stage 47				0.0722456087	15.7575922298
Stage 48				0.0685392852	15.826131515
Stage 49				0.0650231024	15.8911546175
Stage 50				0	15.8911546175

Markov Cohort Analysis Output (Quick) with stage groups expanded

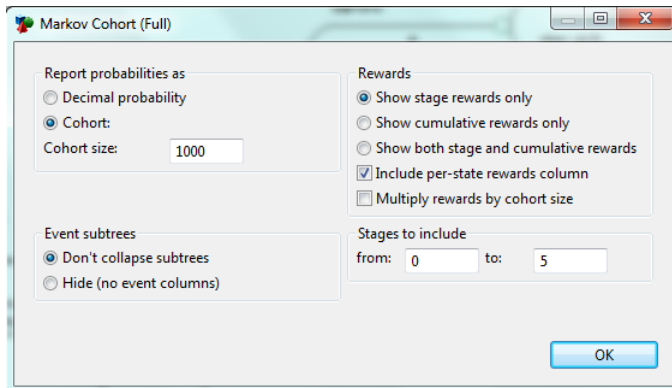
34.3.2 Markov Cohort Expanded Report

If you run the full cohort expected value analysis (not the quick option), the Markov Cohort Analysis output provides an even greater level of detail.

To perform a cohort expected value analysis (full):

- Select the Markov node.
- Choose Analysis > Markov Cohort > Markov Cohort (Full).

- Enter options for generating the cohort analysis output in the Markov Cohort (Full) dialog (see below) and click OK.



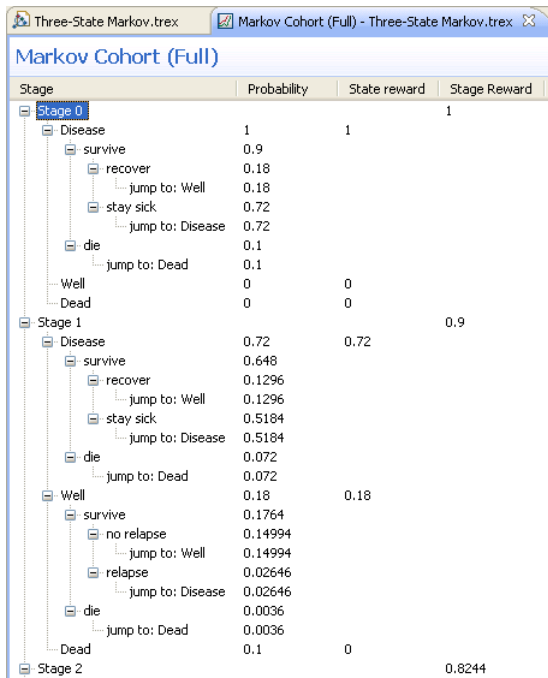
Markov Cohort (Full) Dialog

The options above allow you to customize the output. The options are described below.

- *Report probabilities as*: Determine whether events are displayed using decimal probabilities or as numbers representing probabilities multiplied by an arbitrary cohort size. Rewards are also multiplied by the cohort size.
- *Event subtrees*: Used to summarize probability information. The display of events and their probabilities in the expanded report can be simplified in many models by specifying that similar or cloned events be aggregated ("collapsed"), both within a single state and even across states.
- *Rewards*: Determine which columns of per-cycle and cumulative reward information are reported to the right of the transition information.
- *Stages to include*: Specify which stages to include in the output.

Markov Cohort (Full) Output Options

The expanded output from the "Full" analysis allows you to see the accumulation of rewards in more detail. Specifically, you can see individual transitions within a stage that are not shown in the "Quick" analysis output.



Markov Cohort Output - Expanded

Note that for each stage, all the transitions in all subtrees can be expanded or collapsed. This allows you to check the movement of the cohort through each transition in the analysis. For example, of the 0.9 portion of the cohort that survives, you can see that 0.18 (20%) recovers and 0.72 (80%) stays sick.

34.3.3 Markov Cohort Summary Report

The Markov Cohort Summary Report is a non-grouped grid format for its output. This better matches up with outputs generated from TreeAge Pro 2009 and earlier versions.

Markov Cohort Summary

Stage	% - Disease	% - Well	% - Dead	Reward	Cumulative Reward	Rwd - Disease	Rwd - Well	Rwd - Dead
0	1.000	0.000	0.000	1	1	1	0	0
1	0.720	0.180	0.100	0.9	1.9	0.72	0.18	0
2	0.545	0.280	0.176	0.8244	2.7244	0.54486	0.27954	0
3	0.433	0.331	0.236	0.76432	3.48872	0.43339	0.33093	0
4	0.361	0.354	0.286	0.71437	4.20309	0.36069	0.35368	0
5	0.312	0.360	0.329	0.67122	4.87431	0.31169	0.35954	0
6	0.277	0.356	0.367	0.63286	5.50718	0.27727	0.3556	0
7	0.252	0.346	0.402	0.59803	6.1052	0.2519	0.34612	0
8	0.232	0.334	0.434	0.56591	6.67111	0.23225	0.33366	0
9	0.216	0.320	0.464	0.53601	7.20713	0.21627	0.31975	0
10	0.203	0.305	0.492	0.50799	7.71512	0.20272	0.30528	0
11	0.191	0.291	0.518	0.48161	8.19673	0.19083	0.29078	0
12	0.180	0.277	0.543	0.45672	8.65345	0.18014	0.27657	0
13	0.170	0.263	0.567	0.43317	9.08662	0.17036	0.26281	0
14	0.161	0.250	0.589	0.41088	9.4975	0.16129	0.24959	0
15	0.153	0.237	0.610	0.38976	9.88726	0.15282	0.23694	0
16	0.145	0.225	0.630	0.36974	10.25699	0.14486	0.22488	0

Markov Cohort Summary Report

With no grouping and a standard grid format, this output is often better for exporting to Excel.

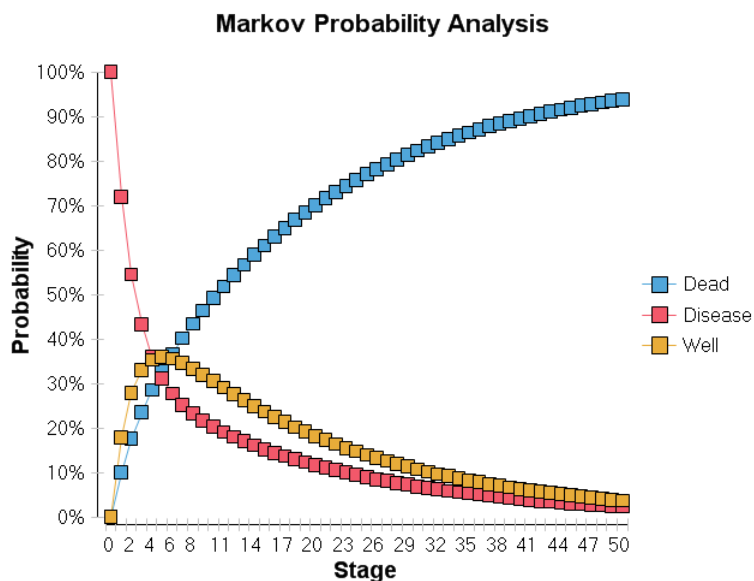
34.3.4 Markov Cohort Graphical Output

You can also generate graphical data from the Markov Cohort Analysis output by clicking on the links to the right of the output data. Each option generates a graph tracking calculated values against time (stage). The options are described below:

- *State probabilities*: This graph plots the changing state probabilities at each cycle, and is closely related to the survival curve.
- *Survival curve*: Survival curves are a standard means of communicating the results of a Markov analysis. TreeAge Pro will prompt you to select which states represent death, and then will plot the sum of the “alive” state probabilities. This graph can also be used to create probability curves that group other kinds of states — to plot disease-free survival, for example.
- *State reward*: This graph shows, for each state, what reward was received at each stage. (For cost-effectiveness models, cost and effectiveness are plotted separately.)
- *Stage reward*: This graph shows the stage reward (sum of all state rewards) accumulated for each stage.
- *Cumulative reward*: This graph shows the cumulative stage reward as it increases with each stage.

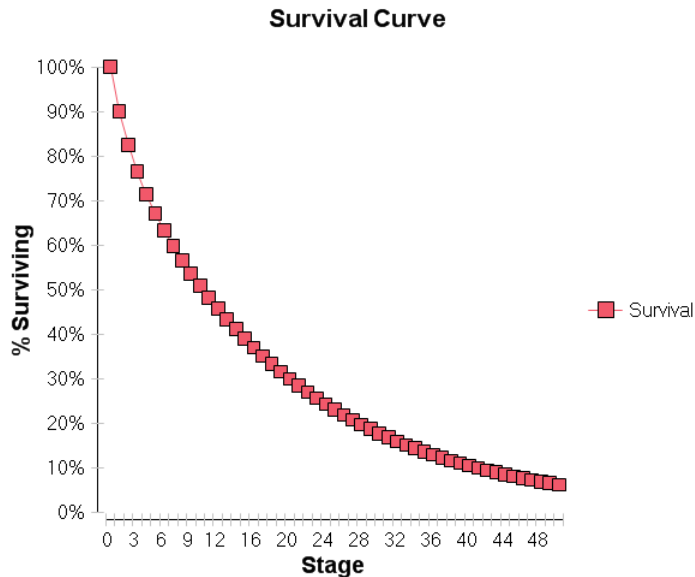
Markov Cohort Analysis graphical output options

The state probabilities graph shows the transition towards the Dead state as time passes.



State Probabilities Graph

The survival curve graph shows the percentage of the cohort that is alive at the beginning of each cycle.



Survival Curve Graph

34.4 A note on microsimulation

Roll back and cohort analysis both evaluate Markov models using cohort calculation methods. Another way to evaluate a Markov model is using Monte Carlo simulation. Microsimulations (a.k.a first-order trials, discrete simulation and individual-level simulation) use random number sequences to send one individual at a time on a single path through the model. This approach is used in some complex models to keep track of an individual's history (beyond what state branch they are in).

Because individuals are randomized based on probabilities, each simulation of a model returns a different set of results, but the summary statistics converge on the true mean as more trials are run. The more complex a model is (e.g. more states, cycles, and small probability events), the more trials required to converge on the expected value.

For a 1-dimensional simulation, where microsimulation is the only loop, the report will include per-individual payoffs/rewards (e.g., life expectancy, cost, etc.) as well as final tracker variable values.

Refer to the Monte Carlo Simulation, Distributions and Probabilistic Sensitivity Analysis Chapter for general information on Monte Carlo simulation. Refer to the Individual-Level Simulation and Markov Models Chapter for more information on specific issues related to building discrete/micro-simulation models.

34.5 A note on half-cycle correction

An important assumption made in discrete-time Markov models is that all state transitions occur simultaneously at the end of each cycle. In reality, however, most kinds of transitions typically occur gradually throughout a time interval (on average, half-way through). This assumption does not affect

reported probabilities (or the survival curve), but it may result in overestimation of expected survival in most models.

The error will be greater if:

- The cycle length is long (e.g., annual).
- There is a significant difference between the rewards associated with the starting state for the cycle and the destination/jump state at the end of the cycle.

Assuming that transitions occur halfway through a cycle on average, the technically ideal correction is to assign a half-reward corresponding to the current state, and then a half-reward at every transition node (transition rewards are described in the next chapter) corresponding to the cost or utility associated with the *jump-to* state.

Most models, however, use a simpler adjustment called the *half-cycle correction*. To implement the half-cycle correction, you can simply divide every alive state's incremental reward in half, and assign the half-reward as its initial and final reward.

In the Three-State Markov example, this will result in a correction of about one-half of a cycle's reward, or about 0.5 years; try the Markov cohort analysis after making this change.

See the next chapter for more on how to use the half-cycle correction.

34.6 Cost-effectiveness Markov models

Building a cost-effectiveness (CE) Markov model is quite similar to working with a regular CE decision tree (refer to the Building and Analyzing Cost-Effectiveness Models Chapter and the Cost-Effectiveness Modeling and Analysis Options Chapter). First, you must set the appropriate preference settings, including calculation method and numeric formatting.

Separate sets of rewards must be entered in the Markov model for the cost and effectiveness attributes. Just as each terminal node in a decision tree can use up to nine payoff expressions, each Markov subtree has nine corresponding reward sets. In CE calculations, reward set #1 might be used for costs and reward set #2 for effectiveness, but this is flexible. If you already have a single-attribute Markov model using payoff set #1 for effectiveness, simply set CE calculations to use payoff set #2 for costs. You also have the option of specifying in the CE preferences that multiple payoffs be combined for cost.

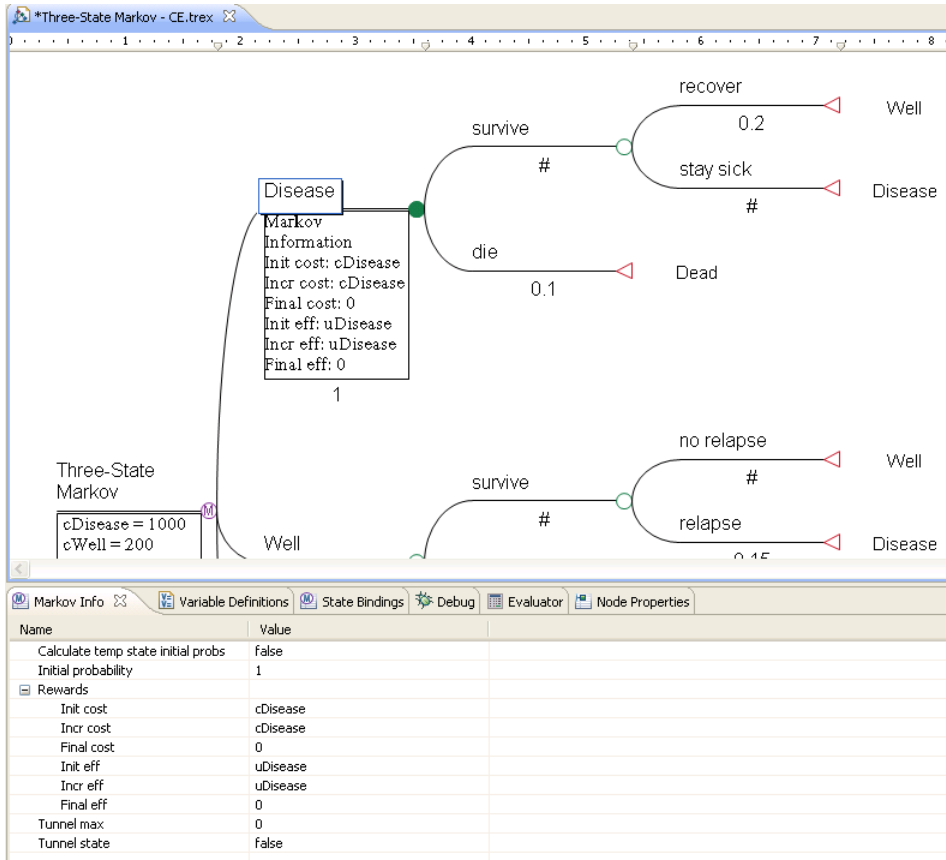
At each Markov state, the three reward types described earlier — initial, incremental and final — can be entered for each reward set.

To assign cost and effectiveness state rewards:

- Select a Markov state.
- Choose Views > Markov Info from the menu.

- In the Markov Info View, enter values or expressions for both cost and effectiveness state rewards.

In a cost-effectiveness tree, rewards are labeled to identify cost and effectiveness rewards in both the Markov Info View and the Tree Diagram Editor.



Cost-effectiveness rewards

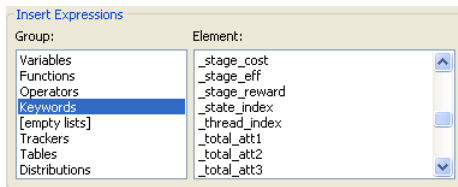
Additionally, a CE Markov model requires that you enter a distinct termination condition for the Cost-Effectiveness calculation method. TreeAge Pro maintains separate termination conditions for each Simple, single-attribute calculation, and for Cost-Effectiveness.

To assign a cost-effectiveness termination condition:

- Select the Markov node.
- Choose Views > Markov Info from the menu.
- In the Markov Info View, enter the termination condition.

34.6.1 Cost-effectiveness keywords

There are several Markov keywords available only in a cost-effectiveness model. The keywords `_stage_cost`, `_stage_eff`, `_total_cost`, and `_total_eff` calculate the single-attribute values; `_stage_reward` and `_total_reward` calculate CE ratios, and are not often used in CE Markov models.



CE keywords

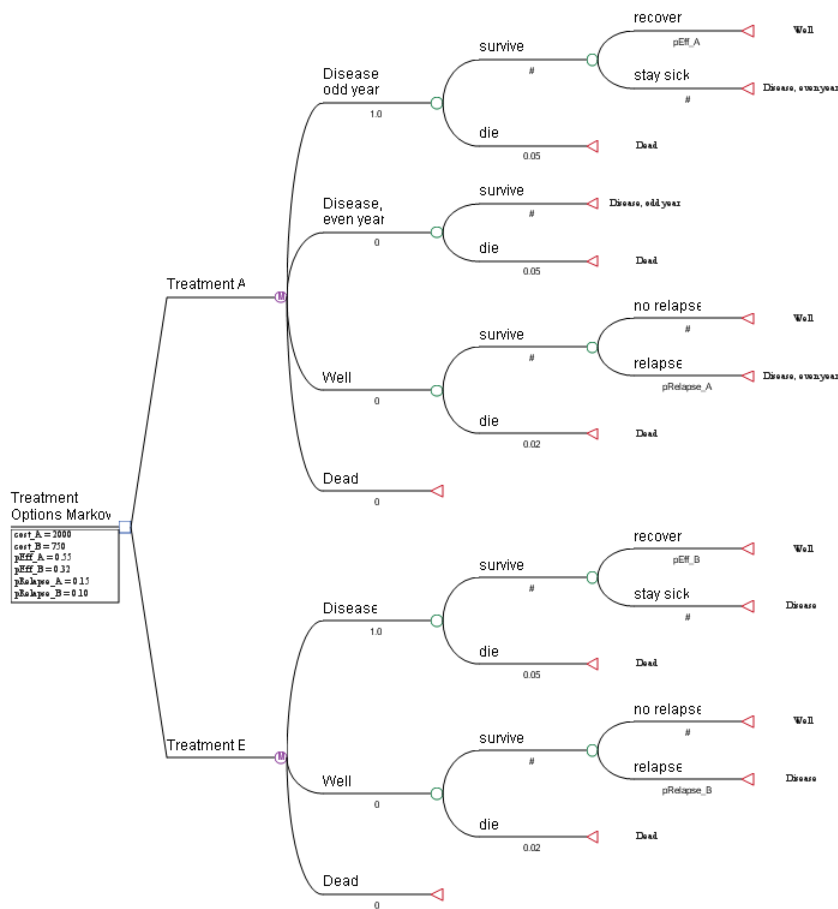
These keywords can be used in expressions within the Markov model. For example, you could create a termination condition `_stage_eff < .001` to stop the analysis when approximately 99.9% of the cohort is dead.



Always change the termination condition to something appropriate for your model! In many models, no effectiveness threshold is needed and the termination condition will only reference the `_stage` keyword. If your effectiveness measure is a rare event counter, then either no effectiveness threshold or a lower one might be required.

34.6.2 A cost-effectiveness Markov model

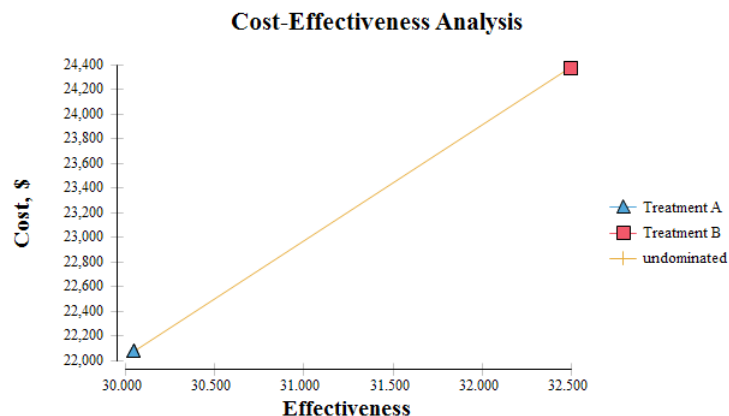
The Health Care tutorial example tree “Treatment Options Markov” combines Markov and cost-effectiveness features. The model compares the cost-effectiveness of two hypothetical treatments using a similar disease model to that of the Three-State Markov process. The tree is shown below .



Treatment Options Markov Tree

In this still relatively simple model, Treatment A is presumed to be faster acting, but cannot be used long term. Treatment B is slower acting, but can be used on a maintenance basis over a prolonged period, effectively preventing more relapses.

You can select one of the Markov nodes at a time and perform a Markov cohort analysis, or select the root, decision node and perform a cost-effectiveness analysis, which will yield the following graph.



Treatment Options Markov Cost-Effectiveness Analysis Results

35. Markov Modeling Tools and Techniques

This chapter covers a number of commonly-used features in Markov models, including using tables of probabilities and discounting rewards. It also covers a number of Markov modeling features that are infrequently used, but may be indispensable in some cases.

Discrete simulation/microsimulation is covered in the Individual-Level Simulation and Markov Models Chapter, and miscellaneous Markov topics are covered in the Markov Technical Details Chapter.

35.1 Keywords, time-dependence, and discounting

TreeAge Pro provides several Markov *keywords* — built-in variables which are available only in a Markov node or its subtree. The first two listed are integer counters:

- `_stage` – the number of cycles that have passed (starts at 0 for first cycle)
- `_tunnel` – the number of cycles spent continuously in a tunnel state
- `_stage_reward` – the reward received by the cohort in the previous cycle (in Simple calculations)
- `_stage_cost`, `_stage_eff` – counterparts of `_stage_reward` in Cost-Effectiveness (CE) calculations
- `_total_reward` – the cumulative reward of all previous cycles; at the end of calculations, this is the overall value of the Markov process
- `_total_cost`, `_total_eff` – counterparts of `_total_reward` in CE calculations
- TreeAge Pro also includes a special function, `StateProb()`, for accessing current state probabilities during analysis.

Markov Keywords

As illustrated in the previous chapter, in the tutorial on building the Three-State Markov model, the `_stage` counter is useful in defining the Markov termination condition. This section will describe other important functions of the `_stage` counter and other keywords.

Refer to the Markov Technical Details Chapter for information on how Markov models are evaluated via the calculation algorithm. This includes when and in what order Markov keyword values are modified, both during cohort analysis and Monte Carlo microsimulation (first-order trials).

35.1.1 Cycle zero

In TreeAge Pro Markov models, the first cycle is referred to as cycle 0 and the `_stage` counter is equal to 0 during this first cycle. For example, if a model's cycle length is one year, cycle 0 represents the first year of the process; if this process started with an individual's birth, cycle 0 would correspond to age 0 – i.e., the year prior to an individual's first birthday.

The following events occur during the first cycle of a Markov process, while the keyword `_stage` is equal to 0:

- The cohort is distributed among the Markov states according to the initial probabilities entered under the branches (the only time these probabilities are used);
- Initial rewards are accumulated based on state membership;
- The members of a state traverse the transition subtree based on the transition probabilities, and the percentage of the cohort at a transition node are assigned the transition rewards in the path back to the state (before entering new states for the next cycle).

You should ensure that references to tables in initial and transition probability expressions, as well as in initial state rewards and transition rewards, will work correctly when `_stage = 0`.

The *incremental* state reward expressions are not accumulated during cycle 0; only the *initial* rewards are evaluated. The initial probabilities determine which states are populated in cycle 0 and where initial state rewards are required. If half-cycle correction is not used, the initial state reward for a state is often the same as the incremental reward; see the section on half-cycle correction later in the chapter for more details.

35.1.2 Using tables of time-dependent transition probabilities – an example

The Three-State Markov model from the previous chapter is an example of a *Markov chain* — a Markov model in which all probabilities and other parameters remain constant over time. In the kinds of Markov models used to represent healthcare issues, however, probabilities and other values often vary over time. This kind of model is referred to sometimes as a Markov process.

In the TreeAge Pro Healthcare module, any expression in a Markov model (not just the termination condition) can reference tables of stage-dependent values using the `_stage` counter. Other kinds of time-dependent expressions can also be created using the `_tunnel` counter, tracker variables, etc.

This tutorial requires two things:

- A copy of the Three-State Markov model you created in the previous chapter. If you did not build the tree, you can open the Health Care tutorial example tree "Three-State Markov".
- A table file called "tMort" to hold time-varying probabilities. Follow the instructions below to create the table. If you have additional questions about working with tables, refer to the Creating and Using Tables Chapter.

To create a new table for use in a tree:

- Choose Views > Tables from the toolbar.
- Click the "add" toolbar button. This will open the Add/Change Table Dialog.
- Enter the table name *tMort* and select the "Use linear interpolation" option for missing rows.
- Click OK to save the table and close the dialog.
- Click the "add" button in the "Table Rows" section of the Tables View. Click seven more times to add a total of eight rows.

- Edit the data in each row to match the data presented below.

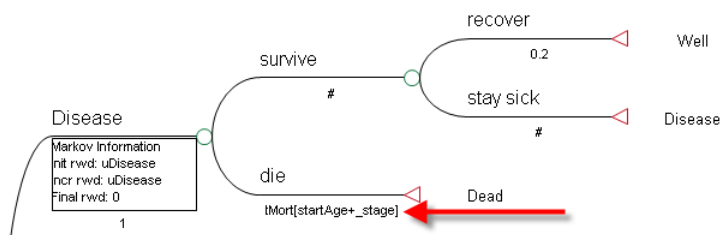
Index	Value1
0.0	0.001
30.0	0.005
40.0	0.007
50.0	0.01
60.0	0.025
70.0	0.05
90.0	0.1
120.0	0.3

tMort table

Now, update the Three-State Markov model to use the new table of mortality probabilities.

To look up a transition probability in a table:

- First define the variable *startAge* at the root node and define it with the value 30.
- Select the *die* branch of the *Disease* state, and change its probability to the formula *tMort[startAge+_stage]*.



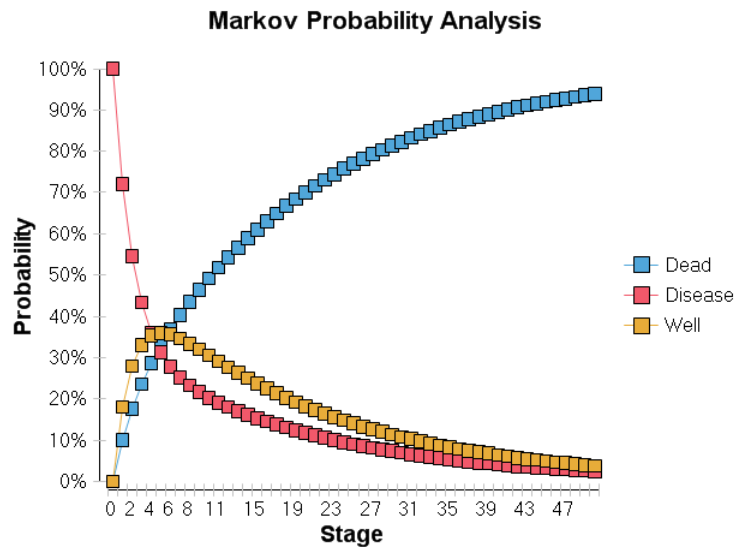
Transition probability from tMort table

Now, in place of a fixed probability of death from other causes, TreeAge Pro will calculate the transition probability at every cycle using the table lookup *tMort[startAge+_stage]*. The first set of transitions in the Markov process, when *_stage* = 0, will use the value returned by the reference *tMort[30+0]*, which is 0.005.

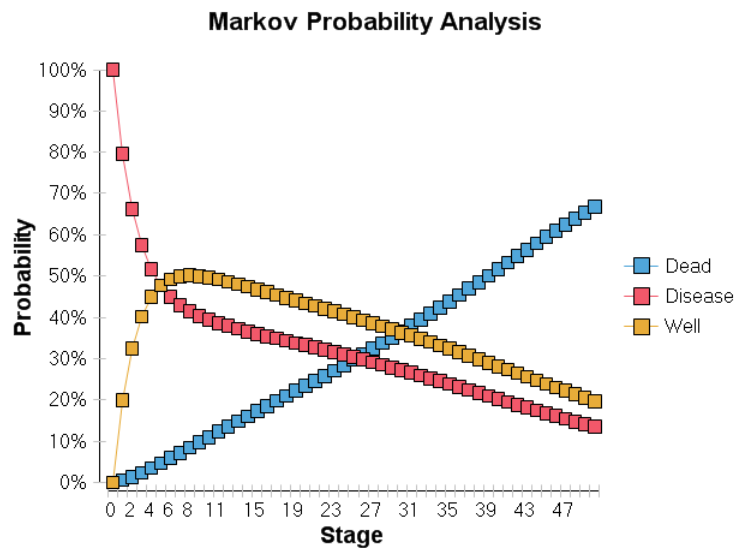
Each subsequent cycle will use a higher mortality probability, because the values in the *tMort* table increase as the indexes increase. Also relevant in this case, when you reference a missing index, the table is currently set to interpolate between existing indexes. In the example, the missing value for cycle 1 when the table reference is *tMort[31]* will be calculated using linear interpolation between the table values for indexes 30 and 40. The interpolated probability will be 0.0052. Missing rows at subsequent cycles will be similarly calculated.

If you now roll back the tree, the Markov node should display an expected value of 34.818 – significantly higher than the roll back value calculated in the previous chapter because the early probabilities of death in the table are lower than the original 0.01 probability of mortality. The updated tree is available with the name *Three-State Markov-Time.trex* for your review.

If you run a Markov cohort analysis in the new version of the tree, and compare the new state probabilities graph with the graph generated in the previous chapter, you will see that the cohort transitions to death considerably more slowly.



State Probabilities Graph - death probability fixed



State Probabilities Graph - death probability from table

35.1.3 Discounting rewards

In addition to defining stage-dependent probabilities, tables can also describe stage-dependent rewards such as costs or utilities. However, if all you need to do is *discount* costs and utilities, a simple exponential formula may be used instead of a table. For example, the expression

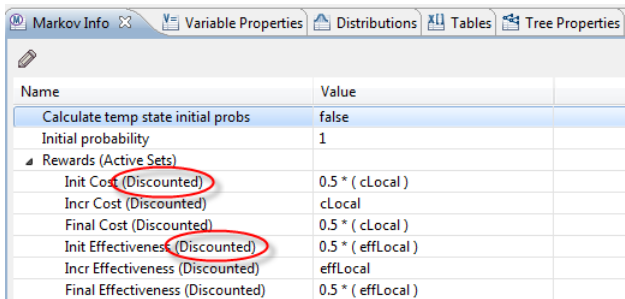
$$\text{cost}X/((1 + \text{rate})^{-\text{stage}})$$

can be used to discount the reward value at each stage. If the cycle length is not equal to the period of the discount rate (usually 1 year), then `_stage` should either be divided (for shorter cycle lengths) or multiplied (for long cycle lengths).

Discounting nearly always needs to be applied to an entire reward set, such as cost and/or effectiveness. TreeAge Pro has two options for discounting.

1. *Apply a single discount rate to an entire reward set:* Use the Tree Preferences to set a specified discount rate to be applied to any given reward set. We call this *Global Discounting*, and we recommend using this approach as it is the easiest to implement.
2. *Adjust each reward manually:* Use the `Discount()` function as described further below to manually change the value of each reward.

To *apply a single discount rate to the entire reward set*, enter the details in the Tree Preferences as described in the Payoff Preferences section. When the Global Discounting is applied to a reward set, the word "Discounted" appears after the reward name in the Markov Info View, as in the figure below.



Name	Value
Calculate temp state initial probs	false
Initial probability	1
▲ Rewards (Active Sets)	
Init Cost (Discounted)	0.5 * (cLocal)
Incr Cost (Discounted)	cLocal
Final Cost (Discounted)	0.5 * (cLocal)
Init Effectiveness (Discounted)	0.5 * (effLocal)
Incr Effectiveness (Discounted)	effLocal
Final Effectiveness (Discounted)	0.5 * (effLocal)

Global Discounting applied to both reward sets for Cost and Effectiveness, indicated by the "Discounted" label after the reward names.

The Healthcare tutorial example model `Markov.Discounting.trex` demonstrates global discounting in a Markov model along with use of the Discount functions for rewards. Note that analysis of the model using each discounting option yields the same results.

If you prefer to manually adjust each individual reward, rather than using TreeAge Pro's Global Discounting function, you can use TreeAge Pro's built-in discounting function, `Discount()`. This `Discount()` function represents the exponential formula above and takes three parameters: *value*, *rate* and *time*. For example, the expression

Discount(costX; rate; _stage)

will yield the same result as the exponential formula shown above. The `Discount()` function is equally applicable to the discounting of costs and utilities.

Name	Value
Calculate temp state initial probs	false
Initial probability	1
Rewards	
Init rwd	Discount(uDisease; 5%; _stage)
Incr rwd	Discount(uDisease; 5%; _stage)
Final rwd	0
Tunnel max	0
Tunnel state	false

Discount function in Markov state reward

TreeAge Pro's built-in functions can be inserted into formulas either by typing them in yourself or by using the Formula Editor or auto-fill.

35.2 Probability/rate conversion functions

The following functions are used to convert between rates and probabilities (or odds).

They generally employ one or both of the arguments *rate* (or *prob*) and *time*. In each case, it is essential that the values for these parameters are based on the same scale. For example, if a rate being converted is in terms of years (such as yearly mortality), the time parameter must also be in years.

Function	Explanation
DEALE(rate; time)	DEALE is an acronym for “declining exponential approximation of life expectancy.” $(1 - e^{-rate*time})/rate$
OddsToProb(odds)	Converts odds into a probability. $odds/(1 + odds)$
ProbFactor(prob; fac)	First converts the probability to odds, then multiplies it by the given factor, then converts it back to a probability. See OddsToProb and ProbToOdds.
ProbToOdds(prob)	Converts a probability into odds. $prob/(1 - prob)$
ProbToProb(prob; multiplier)	Converts a probability into a rate, multiplies the rate by the given multiplier, and converts back to a probability. $RateToProb((ProbToRate(prob; 1) * multiplier); 1)$
ProbToRate(prob; time)	Converts a probability into a rate, and divides the rate by time. $-ln(1 - prob)/time$
RateToProb(rate; time)	Multiplies a rate by time, and converts it into a probability. $1 - e^{-rate*time}$

Probability/rate Conversion Functions

These functions are sensitive to user errors; you are urged to exercise great care when using them in your models. It is recommended that you use the Calculator/Evaluator to test expressions using these functions.

`RateToProb()`, as the name suggests, is used to convert a rate into a probability, either for the same time period (`time=1`) or a different time period. For example, if a disease being modeled has a yearly mortality rate of .05, you could convert this to a probability using the formula:

`RateToProb(.05; 1)`

`ProbToRate()` provides the reverse function to `RateToProb()`, converting a probability for a given time period to a rate. Using the additive property of rates, rates for two events can be added together and converted back to a probability to get the probability of two events both occurring.

In both `RateToProb()` and `ProbToRate()`, the second time parameter will allow you to convert between a rate for an interval of one length and a probability for an interval of a different length.

You may also need to convert a probability for one cycle length to an equivalent probability for a different cycle length. You cannot simply multiply the original probability by a factor. Instead, use the `ProbToProb` function. The following expression changes an annual probability of 0.1 to a monthly probability:

`ProbToProb(0.1; 1/12)`

The `DEALE()` function is cumulative, so the time parameter means “over the course of this amount of time.”

35.3 State/transition probability functions

There are other functions with “Prob” in their names that are not used for conversion between rates, probabilities, or odds. The functions listed here have special applications in Markov models. See the section on Dynamic cohort models later in this chapter for details.

Function	Explanation
<code>StateProb(i)</code>	Returns the state probability of state #i at the start of the current cycle. If running a microsimulation, parallel trials are required or returns as 0. In dynamic models using non-coherent probabilities, effectively returns “counts” instead of true percentages. See note below for more details.
<code>StateProb(_state_index)</code>	Returns the state probability of the state currently being evaluated. The <code>_state_index</code> keyword first returns the index for the current state.
<code>StateProb(i; j)</code>	Returns the sum of the state probabilities of the set of states from #i to #j.
<code>StateIndex("state/binding")</code>	Returns the integer index (starting at 1) of the named state, or the state pointed to by the named binding. Usually used in combination with the <code>StateProb()</code> function. Use in combination with the <code>StateProb()</code> function.

Function	Explanation
	Useful while building the tree, but changing the order of branches or adding branches may affect states' indexes and cause calculation errors.
TransProb()	Returns transition path probability of the current node/branch – i.e., the product of the most recently cached transition probabilities. The product does include the probability stored for the node/branch where the calculation is being called/used. However, note that if branch X's probability is being calculated, then branch X's cached probability from the previous cycle will be used in the product!
PathProb()	Outside of the Markov transition subtree, returns the cumulative path probability for the node being calculated (up to the Markov node in a Markov subtree). Use TransProb() within the Markov subtree.

State/Transition Probability Functions

An argument in StateProb() should correspond to the integer index of a branch of the Markov node (e.g., index=1 for the top state, StateProb(1)).

To return the sum of the state probabilities of a range of states, specify the range of branch indexes using two arguments. For example, use StateProb(2;5) to add the state probabilities of states 2 through 5.

Use TransProb() in combination with StateProb() – and perhaps even PathProb() – to determine the percentage of the population/cohort that experience a particular transition at a particular point in time.

If state #n is a tunnel state, StateProb(n) returns the total state probability of all temporary states. StateProb(-x) returns only temporary state x's state probability.

The StateProb() function will work with non-coherent probabilities in a dynamic cohort model.

35.4 Assigning onetime costs and utilities

There are a number of different situations which may require assigning a *onetime* reward in a Markov model, rather than an incremental reward for each cycle spent in a particular state.

35.4.1 Half-cycle correction

Real processes occur in continuous time, with transitions and other events occurring throughout an interval of time. In TreeAge Pro, however, a Markov process occurs as a discrete sequence of snapshots, with transitions understood to occur at the end of each cycle. As described in the note on half-cycle correction in the previous chapter, in an absorbing Markov process where everyone dies eventually, an uncorrected expected value calculation will overestimate life expectancy by about half of a cycle (0.5 years in a one-year cycle length model). The explanation for this is relatively simple.

In whatever cycle a “member” of the cohort analysis dies, they have already received a full cycle’s worth of state reward, at the beginning of the cycle.

In reality, however, deaths will occur halfway through a cycle on average. So, someone that dies during a cycle should lose half of the reward they received at the beginning of the cycle (e.g., -0.5 years of life expectancy in a one-year cycle length model).

However, instead of implementing the half-cycle correction as a toll at each transition to death, it is easier to implement it in an absorbing process simply by subtracting a half-reward from the rewards assigned at the beginning of the process, in cycle 0 — i.e., by setting a state’s initial reward to one-half of its incremental reward. This is the primary, though not only, reason that the state rewards are separated into three parts.

In a non-absorbing process, in which a significant percentage of the cohort may be alive when the process terminates, cohort members still alive at the end of the process should be given back the half-cycle “death” correction taken from their initial reward at the beginning of the process. This is done by adding on a half-reward after termination in the final reward for all alive states (it does not hurt to always include the initial and final components at every state).

To perform half-cycle correction:

- Select a Markov state node.
- Open the Markov Info View.
- Enter the initial and incremental rewards.
- Select a reward for the appropriate payoff set.
- Click on the "pencil" icon in the view's toolbar to open the State Reward Dialog.
- Click the Half-Cycle Correct button.

The initial and final rewards will be updated.

Node: Disease

Initial Reward, Set #1 =
 $0.5 * (uDisease)$
 Calculated value: 0.5

Incremental Reward, Set #1 =
 uDisease
 Calculated value: 1.0

Final Reward, Set #1 =
 $0.5 * (uDisease)$
 Calculated value: 0.5

Insert Expressions:

Group:	Element:
Variables	
Functions	
Operators	
Keywords	
[empty lists]	
Trackers	
Tables	
Distributions	

Definition info:

Markov State Info
 Description: Disease
 Comment:

OK Cancel

Half-cycle correction via State Reward Dialog

Any reward that is a function of life expectancy (i.e., medication costs that occur gradually over a cycle) is usually corrected in the same way.



In models that calculate quantities other than simple life expectancy, for example *quality-adjusted* life expectancy, different alive states will have different rewards. This means that a perfect half-cycle correction might require correcting not just for death transitions, but for other kinds of transitions from higher value states to lower value states (i.e., where someone should receive half a cycle of the starting state's reward and half of the ending state's reward). Note, however, that Markov approximation errors in two strategies will often cancel each other out in incremental calculations, and reasonable judgement should be used to decide when to use half-cycle correction.

35.4.2 Prior costs

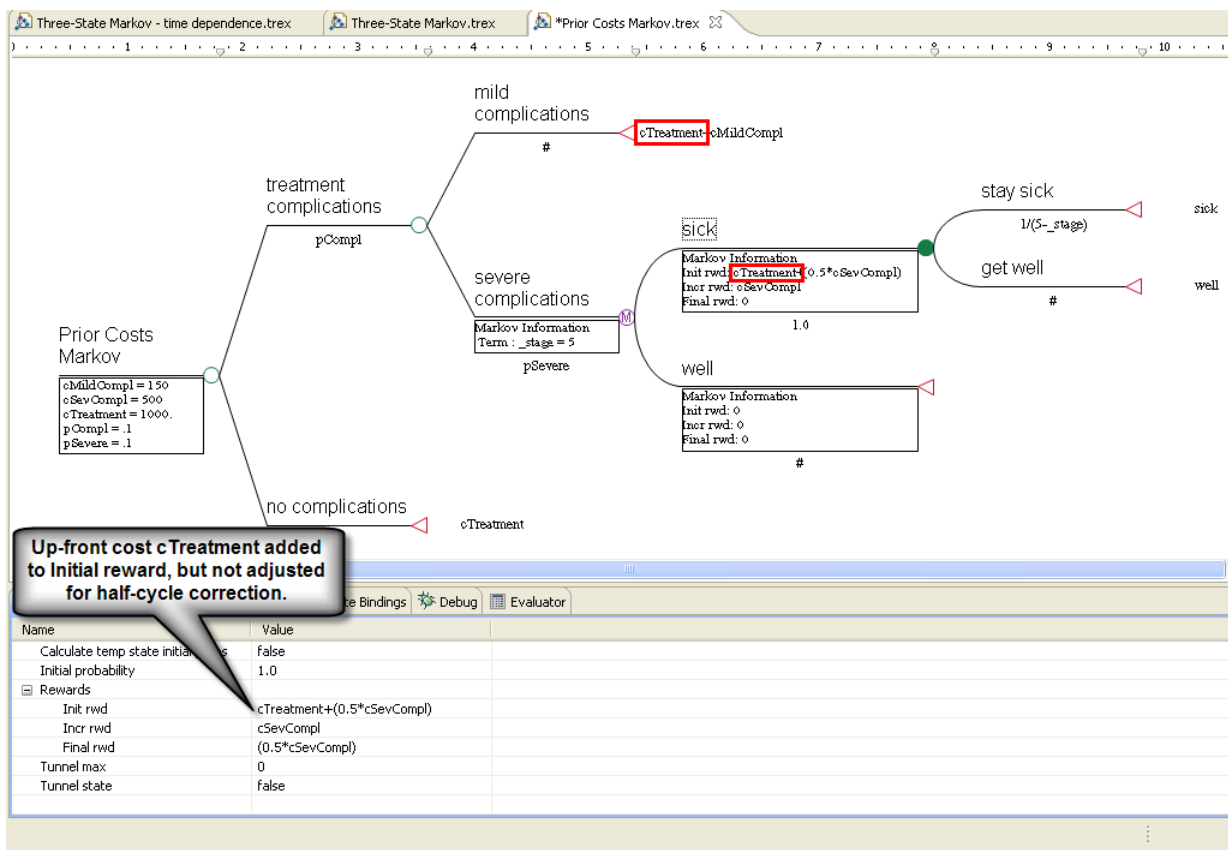
In some models, it is necessary to account for costs, utilities, or life expectancy that occurred prior to the Markov process. Consider, for example, a tree which deals with the uncertainties associated with a particular treatment. In this model, a Markov process will be encountered only if a particular event occurs. In the standard tree structure, costs are incorporated into a payoff formula at terminal nodes. In the scenario including the Markov model, though, these costs must also be accounted for in Markov rewards.

Typically, prior value expressions should be entered in the initial state reward of all states with a nonzero initial probability. This would ensure, for example, that all members of the cohort receive the prior costs. In a cost-effectiveness model (or any model with multiple attributes) be sure to use the appropriate reward set. Cost-effectiveness Markov models are discussed later in this chapter. If your model also uses the half-cycle correction (see above), the initial reward expressions must combine the prior values and the half reward.

To include prior costs in a Markov model:

- Create and define a variable or expression that represents all costs accumulated before the Markov process.
- For each state with a nonzero initial probability, update the initial reward expression to add the prior costs expression, remembering to keep the half-cycle correction if needed.

The Health Care tutorial example tree "Prior Costs Markov" is shown below.



Prior Costs Markov Model

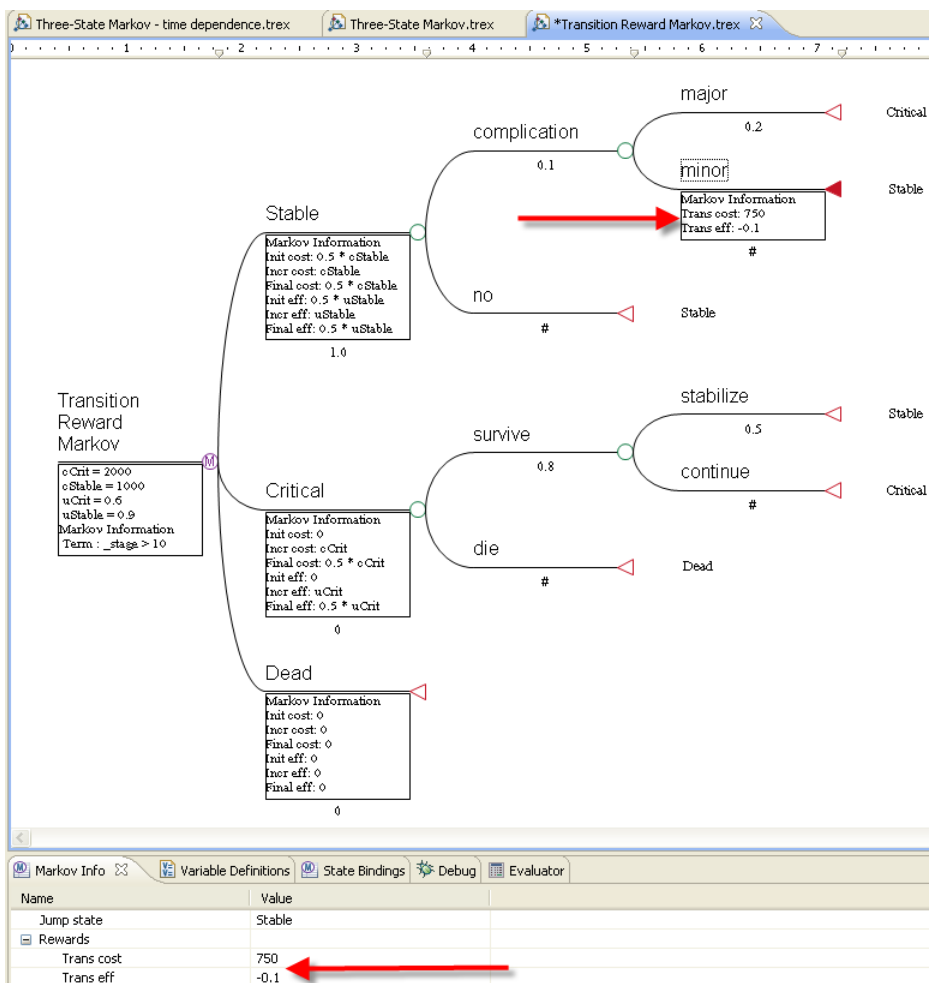
35.4.3 Transition rewards

In some models, you may need to account for a cost or disutility associated with a transient event rather than a state. In many such cases, a transition reward can be used. Transition rewards can be assigned at any node to the right of the Markov state nodes (not just the actual transition nodes).

For instance, a onetime cost may be associated with admission as an inpatient. This cost is not incremental and should not be accumulated in each interval spent in the hospital. Nor can the cost be assigned using an initial state reward if the admission event is not just an initial, cycle 0 event (initial rewards are only assigned when `_stage = 0`).

Another example might be a relatively minor complication event during treatment. Although the complication is not a state itself, and may have no effect on state transition, it may have costs and/or disutilities associated with it.

In the Health Care tutorial example tree "Transition Reward Markov", transition rewards are specified both for costs (in reward set #1) and for effectiveness (in reward set #2).



Markov Transition Reward Tree

To assign a transition reward:

- Select the node where the event occurs, to the right of a Markov state.
- Choose Views > Markov Info from the toolbar.
- Enter values in one or more of the fields Rewards > Trans <Reward Type> in the Markov Info View.

If the Markov information display preference is turned on, transition reward expressions are shown below the branch.

Notes on transition rewards:

- Transition rewards are not accounted for separately from state rewards.
- Transition rewards are associated with the *cycle* in which they occur.
- In cohort analysis reports, transition rewards are not associated with the *state* in which they occur; instead, they are divided among the Markov states to which the transition may lead based on the relative transition probabilities.
- Like state rewards, transition rewards are *added* to the net reward. Thus, transition rewards should be entered using the appropriate sign, positive or negative. For example, transition costs are normally entered as positive numbers, while transition disutilities are normally negative numbers.

35.5 Cloning Markov models

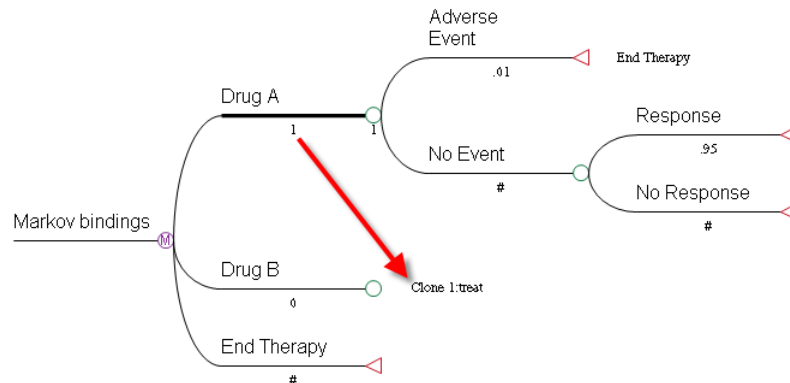
A major advantage of using clones in any tree is the ability to reuse a particular structure in multiple parts of a tree. By doing this, you're still retaining the option to vary probability and other value expressions in each "copy," but you only have to maintain the master subtree (refer to the Tools and Functions for Complex Trees Chapter). In TreeAge Pro, entire Markov models can be cloned within a single tree and parts within a complex transition subtrees can also be cloned.

35.5.1 Using Markov state bindings

Clone copies of Markov transition subtrees will, by default, employ the jump-to state settings specified in the clone master. In TreeAge Pro, Markov bindings can be used to have a transition in a clone copy use a different jump-to state than the clone master.

As described in the Tools and Functions for Complex Trees Chapter, variables can be used in clone copies when numeric values such as probabilities should not be controlled by the clone master. Markov state bindings function similarly; rather than assigning a numeric value to the binding name, a Markov state name is assigned instead. Like variable definitions, Markov state bindings must be defined at an appropriate node. Markov state bindings may be defined at any node on the Markov subtree, including the Markov node. Typically, they are created at the root nodes of the clone master and clone copies.

The use of Markov state bindings can be illustrated using the Health Care tutorial example tree "Markov Bindings", shown below unfinished.



Markov Bindings Tree

Transitions must be assigned to the Response and No Response nodes in the clone master. The Response transition node in Drug A's subtree should jump back to Drug A, while the same node in Drug B's clone copy subtree should, instead, jump to Drug B. Similarly, Drug A's No Response node should point to Drug B, while Drug B's No Response node should point to End Therapy.

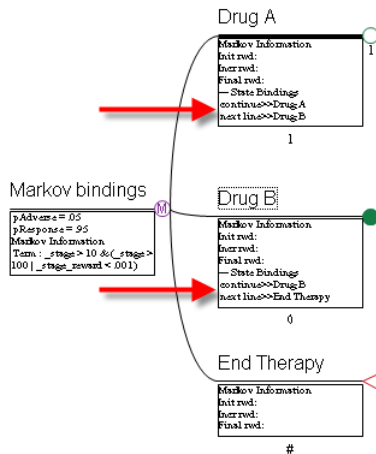
To define Markov state bindings in the Markov subtree:

- Open the Markov Bindings tree.
- Select the *Drug A* node.
- Choose Views > State Bindings from the toolbar.
- In the State Bindings View, click the "add" toolbar icon to add a new binding.
- Enter the Name *continue* and select the State *Drug A*.
- Add another binding and enter the Name *next line* and select the State *Drug B*.
- Select the *Drug B* node.
- In the State Bindings View, add two bindings with the Name/State combinations *continue/Drug B* and *next line/End Therapy*.

Variable Definitions		State Bindings	
		<div> <div>×</div> <div>+</div> </div>	
Name	State		
continue	Drug A		
next line	Drug B		

Create Markov Bindings

If the display of Markov information is turned on in the tree (in the Tree Preferences dialog, under the Variables Display category), Markov bindings will be displayed below other Markov information, in the form *binding name >> jump-to state name*.



Markov Bindings shown in tree

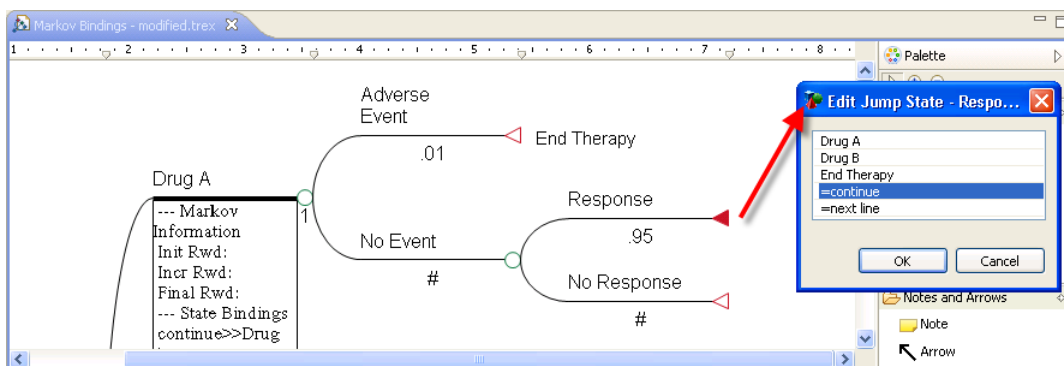
Bindings that you create have no effect until a binding name is referenced at a Markov transition node in a clone master.

When Markov state bindings are found in the path back to the Markov node, the list of options under Jump state in the Markov Info View will include Markov bindings as well as the regular state names. The binding names displayed in the list are prefixed with the equal sign (=) to distinguish them from actual states. (Thus, when naming states, you should avoid using a leading =, although this is not strictly forbidden.)

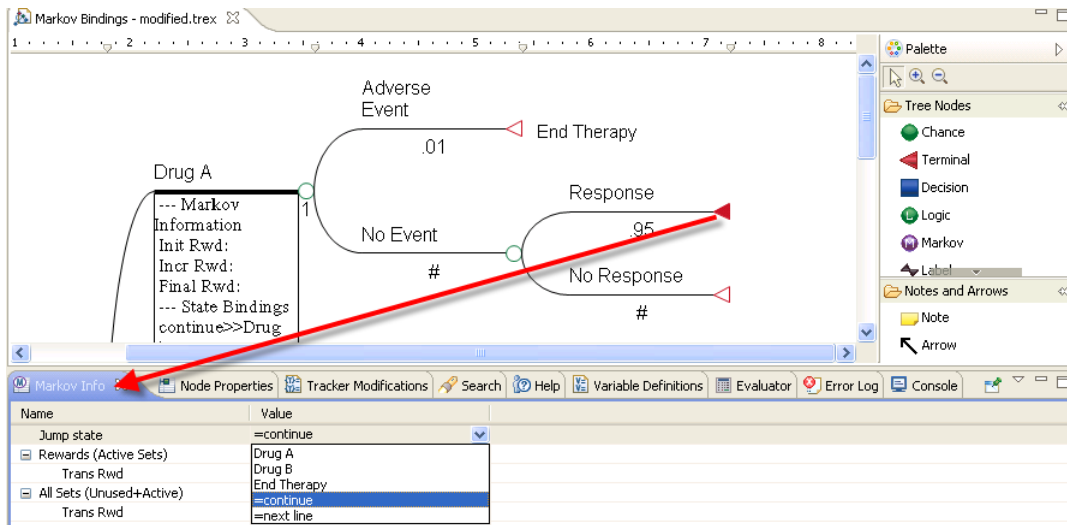
To use a Markov state binding in a clone master transition:

- Double-click on *Drug A's Response* node to open the Edit Jump State Dialog.
- In the Edit Jump State Dialog, select the Markov Binding `=continue` and click OK.
- ... OR...
- Select *Drug A's Response* node.
- Choose Views > Markov Info from the toolbar.
- In the Markov Info View, select the Markov Binding `=continue` for the Jump state.

Repeat these steps to set the jump state for the No Response Node to `=next line`.



Assign Markov binding via the Edit Jump State Dialog



Assign Markov binding via the Markov Info View

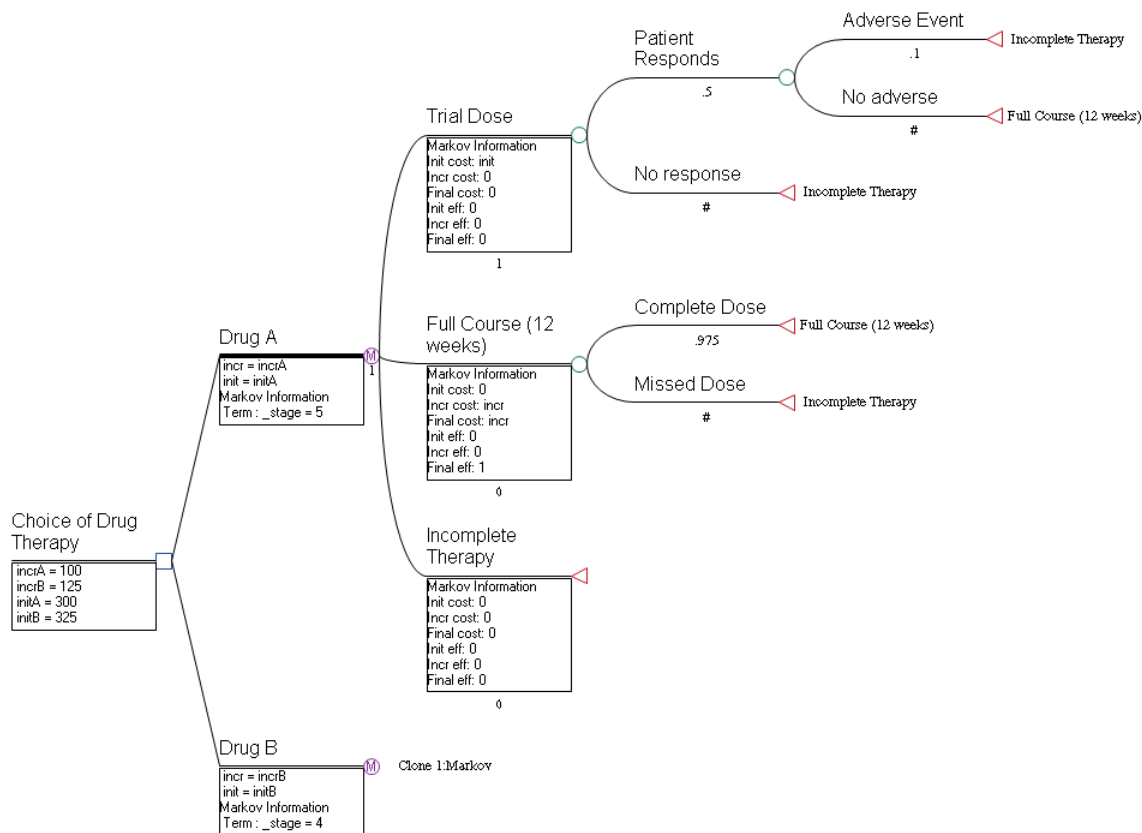
When a binding name is used at a transition node, the search for the binding proceeds in right-to-left fashion, as with variable definitions. Since the Markov state bindings *=continue* and *=next line* are defined differently at the *Drug A* and *Drug B* nodes, the jump states will be different even though the transition subtree is cloned.



To avoid Markov structure errors due to subtle differences in state names, you should use caution when trying to clone part of one Markov process in order to attach copies in a different Markov process. (If you are determined to try, however, TreeAge Pro will allow it!) You might instead try cloning the entire Markov process, creating a clone master at the Markov node. However, there should be no problems with attaching clone copies from one state onto another state within the same Markov process.

35.5.2 Cloning an entire Markov process

Consider the Health Care tutorial example tree "Complex Markov Cloned", shown below.



Complex Markov Cloned Tree

Drug B's transition subtree is a clone copy of *Drug A's* subtree. On the surface, these subtrees appear to be identical, but in fact, the strategies have different termination conditions (assigned at the Markov node, outside the clone master).

State rewards utilize the variables *init* and *incr*. These variables are defined differently at the two Markov nodes resulting in different cost calculations. Similarly, different values could be used for each subtree's probabilities simply by converting the numeric probabilities to variables in the clone master, and then uniquely defining the variables at both the *Drug A* and *Drug B* Markov nodes.

35.6 Counting "time in state" with tunnels

A tunnel state can be used when you need to keep track of the number of cycles an individual has remained in a particular state. In a cancer state, for example, transition probabilities to other states often depend on how long the individual has been in the cancer state.



Tunnels versus simulation/trackers:

Instead of using tunnel states, some Markov models are built using *tracker variables* and *microsimulation* to count time-in-state; each approach has its advantages. Tunnel states work during both cohort analysis (i.e., rollback, n-way sensitivity analysis) and microsimulation. Trackers are more restrictive, *requiring* microsimulation (a.k.a. individual-level simulation) which is generally a more time-consuming analysis. Trackers, however, are extremely flexible, and can be used to keep track of unlimited continuous and discrete states, transient events, etc. Models built using tracker variables can be made structurally simpler, with fewer states/branches. Refer to the Individual-Level Simulation and Markov Models Chapter for details on tracker variables.

If a model is already using trackers/microsimulation, then it will be more efficient/parsimonious to track time-in-state, rather than “tunnel” it.

35.6.1 Temporary states and the `_tunnel` counter

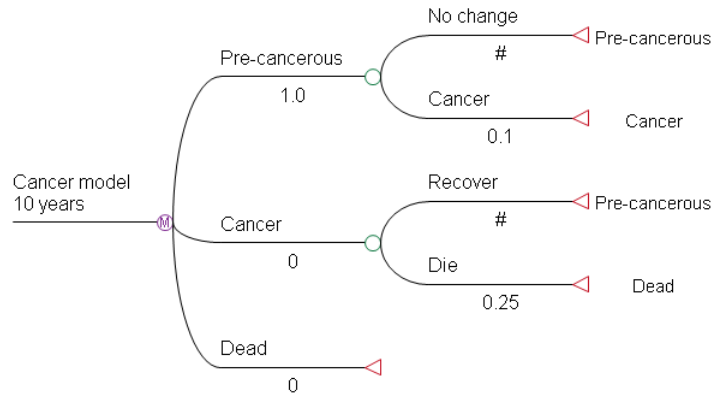
In the Markov modeling literature, a *temporary state* is a state which an individual must exit after one cycle and a *tunnel* is a series of temporary states. Normally, an individual entering the tunnel state — either from another state, or at the start of the Markov process — enters temporary state #1. If an individual remains in the tunnel state for another cycle, they move in order through temporary states #2, #3, and so on.

One way to model a tunnel is to use a separate state for each temporary state, and manually set up ordered transitions between the states. However, TreeAge Pro allows you to represent a tunnel more efficiently, using a single branch from the Markov node; this makes it easy to create tunnels of any length (even with thousands of temporary states).

When you create a tunnel state in TreeAge Pro, all temporary states will use the same transition subtree. In order to specify different transitions for particular temporary states, you can refer to TreeAge Pro’s temporary state counter, a Markov keyword called `_tunnel` (similar to `_stage`). TreeAge Pro starts the `_tunnel` counter at 1 for someone entering a `_tunnel` state, and increments the counter by 1 each cycle they remain in the tunnel state. Using `_tunnel` to count “time in state,” your transition probabilities can look up appropriate values from tables. Logic nodes and statements can also use the `_tunnel` counter.

35.6.2 Using a tunnel state – an example

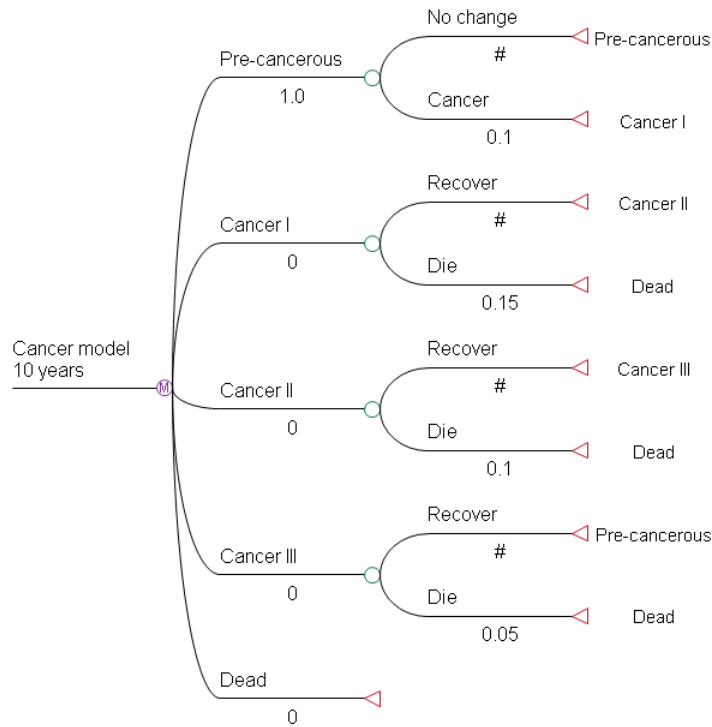
Consider the three-state cancer model shown below. Note that since there is no transition from Cancer back to itself, the patient spends one cycle in the Cancer state before exiting to another state, and therefore Cancer is a temporary state (although not a tunnel yet).



Cancer model before tunnel

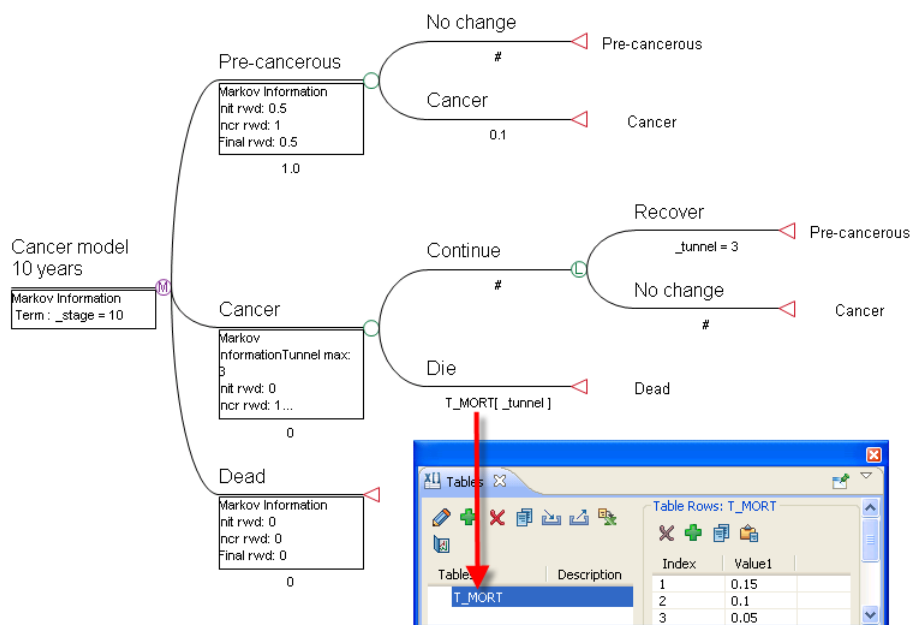
This model may not accurately represent the basic process of the disease. Cancer should probably unfold in a series of temporary states, with different probabilities of changing state (i.e., remission or death) in each successive cycle/year. And these probabilities should depend on how many cycles someone has spent in the Cancer state (which is not given by the `_stage` counter, since everyone starts in Pre-cancerous, and a transition to Cancer may occur at any cycle).

An “exploded” version of the cancer Markov model is shown below. An explicit chain of temporary states is used to describe each year of the cancer. While this more detailed model is still relatively small, as the required number of temporary states increases, explicitly representing all of them becomes more problematic. In TreeAge Pro, the chain of temporary states can be represented more efficiently using a single tunnel state.



Cancer model "exploded" with temporary states

The same Markov process (i.e., calculating the same results) can be built in TreeAge Pro with Cancer set as a tunnel state, as illustrated below. See the Health Care tutorial example tree "Cancer Tunnel".

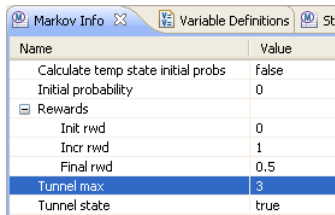


Cancer Tunnel Model

To change a state to a tunnel state:

- Select the appropriate Markov state.

- Choose Views > Markov Info from the toolbar.
- In the Markov Info View, set the Tunnel Max value equal to the maximum required number of temporary states.



Name	Value
Calculate temp state initial probs	false
Initial probability	0
Rewards	
Init rwd	0
Incr rwd	1
Final rwd	0.5
Tunnel max	3
Tunnel state	true

Tunnel Max entry in Markov Info View

The number specified determines how high the `_tunnel` counter will increment (corresponding to the number of copies of the state which TreeAge Pro keeps track of internally during calculations). Individuals that reach the last temporary state and transition into the state again will simply remain in the last temporary state.

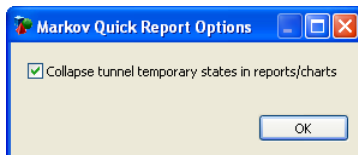
Set the number of temporary states to the minimum number that will account for all differences among the temporary states. For example, if the only difference among the temporary states is a probability that changes for the first four cycles then stabilizes, you will need four temporary states.

35.6.3 Using the `_tunnel` counter

Note the use of the `_tunnel` counter/keyword in probabilities in the Cancer transition subtree shown in the prior section. The *Die* node probability is `T_MORT[_tunnel]`. As a portion of the cohort remains in the state, `_tunnel` increases, and a different probability is pulled from the table.

35.6.4 Merging results for temporary states

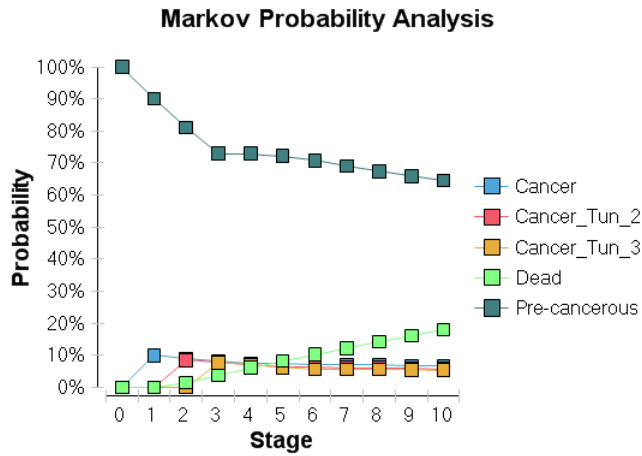
When you perform a Markov cohort analysis at a Markov node that includes a tunnel state, you are given the option to merge each tunnel's temporary states into a single text report column or graph line. In fact, that is the default presentation of data.



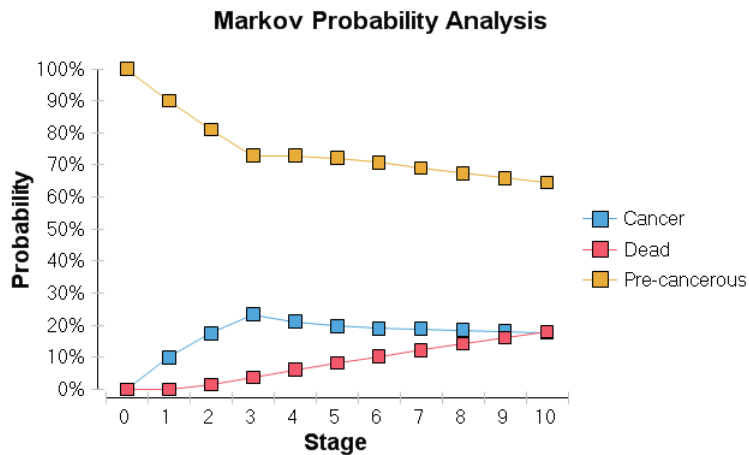
Collapse Tunnels Dialog

This is particularly useful if there are many temporary states in a tunnel. Prior to running the analysis, TreeAge Pro will present the prompt shown above, asking whether or not to merge temporary states. If you choose not to merge temporary states, the text report will append extra columns for the second and subsequent temporary states.

One positive result of merging a tunnel's temporary states is a simpler, more coherent line graph, as shown below.



State Probabilities Graph with Tunnels expanded



State Probabilities Graph with Tunnels collapsed

35.6.5 Populating temporary states at cycle 0

Normally, the initial probability expression assigned to a tunnel state is used to populate only the first temporary state. It is possible, however, to distribute members of the cohort among the different temporary states at cycle 0. The Tunnel Info section of the Markov State Information dialog includes an advanced setting that will cause TreeAge Pro to evaluate the initial probability expression in a tunnel state for every temporary state.

Name	Value
Calculate temp state initial probs	true
Initial probability	.25
Rewards	
Init rwd	0
Incr rwd	1
Final rwd	0.5
Tunnel max	4
Tunnel state	true

Set initial probabilities for tunnel states

In the above example, 25% of the cohort should start the analysis in each of the 4 temporary states associated with this tunnel state.

This could be used, for example, to create a model that uses the `_tunnel` counter to track the age of the members of the cohort with a realistic age distribution. Every state could be made a tunnel state, with their initial probabilities referencing tables of probabilities using the `_tunnel` counter (creating the age distribution).

35.6.6 Special binding names and tunnels

Markov bindings were first discussed earlier in this chapter within the context of clones.

TreeAge Pro supports three special Markov binding names:

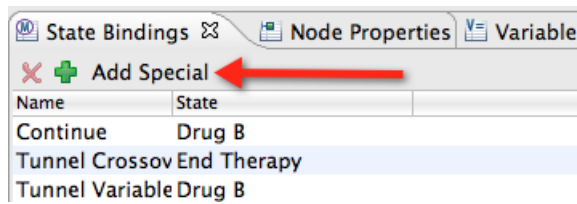
- Tunnel Crossover
- Tunnel Variable
- Tunnel Table

The *Tunnel Crossover* binding is used at a transition node to allow the `_tunnel` counter to increment uninterrupted when transitioning from one state to a different state, provided both are tunnel states. Normally, the `_tunnel` counter would reset to 1 when moving to a different state. If the tunnel crossover value exceeds the maximum number of tunnels for the state, the maximum tunnel value is used.

The *Tunnel Variable* binding is used to dynamically point any transition node to a specific temporary state in the destination tunnel state. The variable must have the name "tunnel_" appended to the front of the starting health state (replace spaces with underscores). See example model Markov Tunnel Bindings for details.

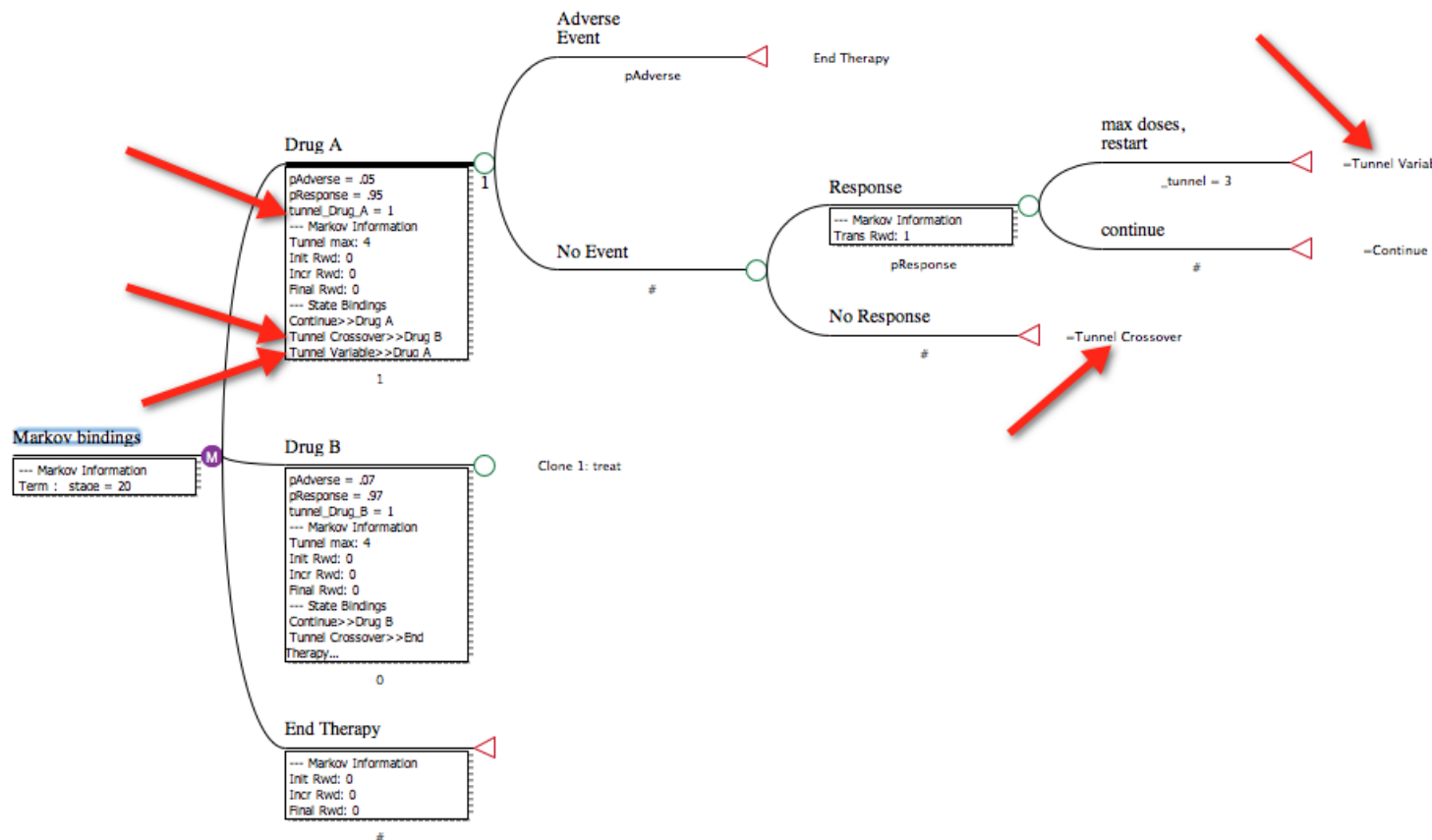
The *Tunnel Table* binding is used in a similar way to Tunnel Variable, except that transitions to particular temporary states are determined by numbers pulled from a column in a TreeAge Pro table. The tunnel table binding will have a name like Tunnel Table C1 where C1 refers to the first value column (not index column) from a table with the name "tunnel_" appended to the front of the starting health state (replace spaces with underscores). When the cohort reaches the tunnel table binding, it looks for the table, finds the row with an index equal to the current `_tunnel` value, then goes to the appropriate column (C1, C2, etc.) to determine the `_tunnel` value for the next cycle, regardless of which jump state is selected for the binding. See example model Markov Tunnel Bindings Table for details.

To create a special binding, click "Add Special" on the State Bindings View toolbar. See below.



Create special bindings (Mac)

The Health Care tutorial example model "Markov Tunnel Bindings" illustrates the use of the Tunnel Crossover and Tunnel Variable special bindings.



Special Bindings (Mac)

The Tunnel Variable binding definition made at the *Drug A* node points to the *Drug A* state. The use of this binding at the *max doses, restart* node within the *Drug A* transition subtree means that this transition will return to the *Drug A* state for the next cycle. However since the binding is a Tunnel Variable, TreeAge Pro searches for variable *tunnel_Drug_A* ("tunnel_" + <state name> replacing spaces with underscores) to determine the temporary state for the next cycle. In this case the variable *tunnel_Drug_A* is defined as 1, so the temporary state will be 1 for the next cycle (restart the doses).

The Tunnel Crossover binding definition at the *Drug A* node points to the *Drug B* state and vice versa. This allows the transition for the *No Response* node to transition to the other state. However, since it is a Tunnel Crossover binding, the temporary state is maintained. The flow would pass from temporary state 1 in the *Drug A* state to temporary state 2 in the *Drug B* state and vice versa.

35.7 Markov decision processes

Markov decision processes are not currently supported.



We strongly recommend having a single decision node in healthcare models. This is the only way that all strategies can be shown in analysis results for comparison.

If there is a downstream decision, it is best to predefine all decisions in a single strategy. For example, if you have a choice of treatment A or B this year and then treatment C or D next year, four strategies could be *A then C*, *A then D*, *B then C* and *B then D*.

35.8 Dynamic cohort models

This section deals with dynamic “cohort” models. Using parallel trails, dynamic populations can also be modeled in microsimulation trials (refer to the Individual-Level Simulation and Markov Models Chapter for details).

35.8.1 Budget impact and infectious disease models

In decision analysis or cost-effectiveness analysis, a Markov node can be used to calculate expected (i.e., mean) cost and effectiveness for a closed population/subgroup of *indeterminate size*. In a budget impact analysis, however, the goals are different; a modified approach to the Markov cohort may be used, including modeling an open (i.e., dynamic) population, implying a *determinate* initial size.

For additional background on budget impact modeling, refer to:

http://www.ispor.org/workpaper/budget_impact.asp

Another context in which a dynamic cohort is relevant is in modeling infectious disease, when force of infection, herd immunity, and other possible factors may depend in part on the determinate *size* of the infectious and/or susceptible population at any particular time point.

35.8.2 Dynamic “populations” and non-coherent probabilities

It has been noted numerous times in the tutorials that TreeAge Pro requires that the branch probabilities of each chance and Markov node in your tree always sum to 1.0. Probabilities that meet this requirement are referred to as “*coherent*.” In certain situations it is useful to be able to remove the probability coherence requirement. This is the case with handling dynamic Markov cohorts.

A *non-coherent* approach to initial and transition probabilities can be used to dynamically resize the cohort at any stage of the analysis. (The approach in a microsimulation model is almost identical; see next chapter.)

TreeAge Pro includes a preference to turn off the coherence error checking that normally protects against the assignment of bad probabilities. This can be done either for the entire tree or for specific nodes. It is recommended that node-level probability overrides are used, so that bad probabilities at other nodes throw the appropriate errors.

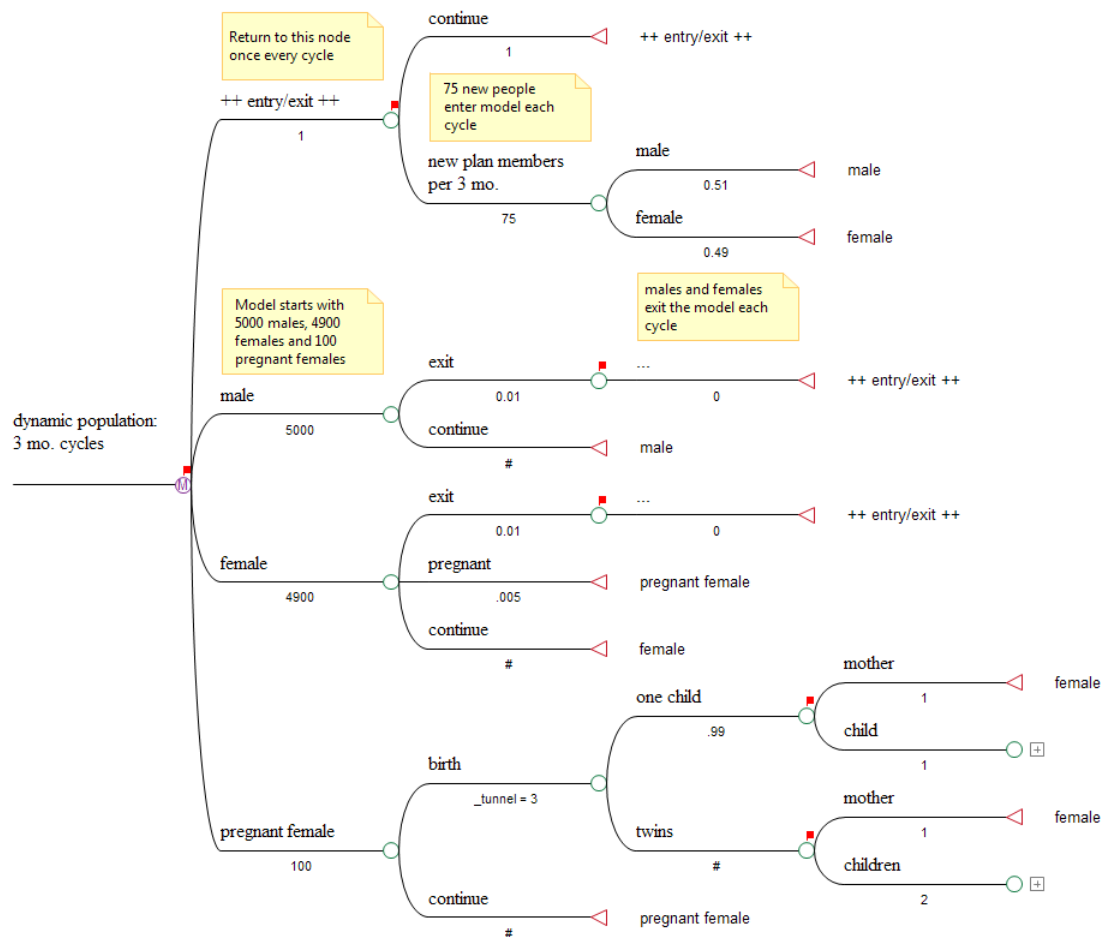
For instructions on how to allow non-coherent probabilities, click one of the following links.

- Allow non-coherent probabilities for specific nodes.
- Allow non-coherent probabilities for the entire tree.

The Health Care tutorial example tree – “Dynamic Population v2008” – is shown below. This model illustrates two key aspects of the use of non-coherent Markov probabilities to model a dynamic/open population:

- Discrete initial sizing of the cohort (e.g., N=10000) using non-coherent *initial* probabilities
- Population growth (e.g., through births) using non-coherent *transition* probabilities

(Note that using only the first option does not really model a dynamic/open population, but rather multiplies results by the initial size.)



Dynamic Population v2008 Model

In the Dynamic Population example, population growth is modeled (the context might be a health care network). The starting population is specified with numbers of individuals in the initial state probabilities — a total of 10000, ignoring ++ *entry/exit* ++. During 10 years' worth of three-month cycles, population growth occurs in two ways: A) entry from other populations, i.e. plan enrollments; and B) births and other changes in family size.

The ++ *entry/exit* ++ state is set up to always have a state probability of 1 (both via its initial probability and via the transition probability for node *continue* returning 1 to that state). This makes its job — injecting new cohort members every cycle — easy. Its state probability will be multiplied by the non-coherent transition probability for node *new plan members per 3 mo*, resulting in an increment of 75 split between the state “probabilities” of the *male* and *female* states.

Births to plan members also use non-coherent transition probabilities (1 mother + 1.01 infants).

Finally, population loss is modeled: note the 0 probability transitions, which subtracts the arriving portion of the cohort (not sent to any state).



In some models, just setting a size on the initial cohort (i.e., using number of individuals starting in each state, instead of initial probabilities) might be useful, perhaps in combination with the `StateProb()` function described earlier. A state's "probability" would actually be a number of "individuals". In an infectious disease model, the probability of a new infection could depend on the number in infected states, for example:

$$p_{Inf} = .2 * \text{StateProb}(6;10) / \text{StateProb}(1;10)$$


Dynamic simulation models:

Non-coherent probabilities are compatible with Markov microsimulation when parallel trials are used. The example model used here is also used to demonstrate dynamic parallel trials simulations as described in the next chapter.

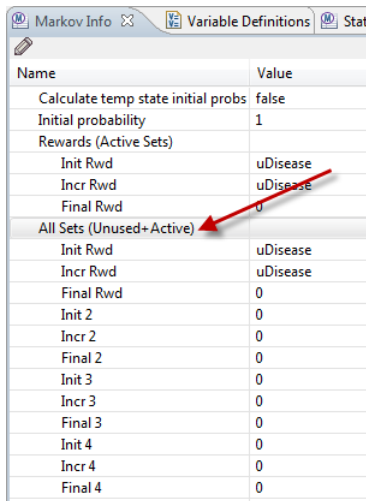
35.8.3 Notes on dynamic-sized cohorts

- Non-coherent probabilities *are compatible* with Markov microsimulation models. The preference described above is used in combination with the "parallel trials" microsimulation option. Refer to the next chapter for details on requirements.
- As a precaution against unintended use, the status bar at the bottom of the TreeAge Pro window displays text to notify you that this setting is active (refer to the Advanced Chance Node Tools and Techniques Chapter).
- Instead of having TreeAge randomize whole individuals as described above, you can selectively and carefully use the rounding functions (Floor, Ceiling, or Round) to the same effect.
- If non-coherent probabilities aren't used to model a finite-sized population, but instead to enable non-exclusive chance node branches — i.e., to create the possibility of going down more than one path — care must be taken not to double-count payoffs.

35.9 Extra Rewards

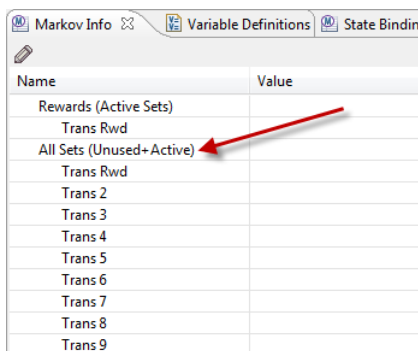
The Tree Calculation Methods and Preferences Chapter described how to use extra payoffs for additional calculations beyond the active payoffs for the specified calculation method. You can enter state and transition rewards for extra payoffs in a Markov model as well.

For Markov state and transition nodes, you can enter rewards for additional payoff/reward sets in the Markov Info View. All enabled payoff/reward sets are displayed beneath the active reward sets. See below.



Name	Value
Calculate temp state initial probs	false
Initial probability	1
Rewards (Active Sets)	
Init Rwd	uDisease
Incr Rwd	uDisease
Final Rwd	0
All Sets (Unused+Active)	
Init Rwd	uDisease
Incr Rwd	uDisease
Final Rwd	0
Init 2	0
Incr 2	0
Final 2	0
Init 3	0
Incr 3	0
Final 3	0
Init 4	0
Incr 4	0
Final 4	0

Extra reward sets in Markov Info View - Markov state node



Name	Value
Rewards (Active Sets)	
Trans Rwd	
All Sets (Unused+Active)	
Trans Rwd	
Trans 2	
Trans 3	
Trans 4	
Trans 5	
Trans 6	
Trans 7	
Trans 8	
Trans 9	

Extra reward sets in Markov Info View - Markov transition node

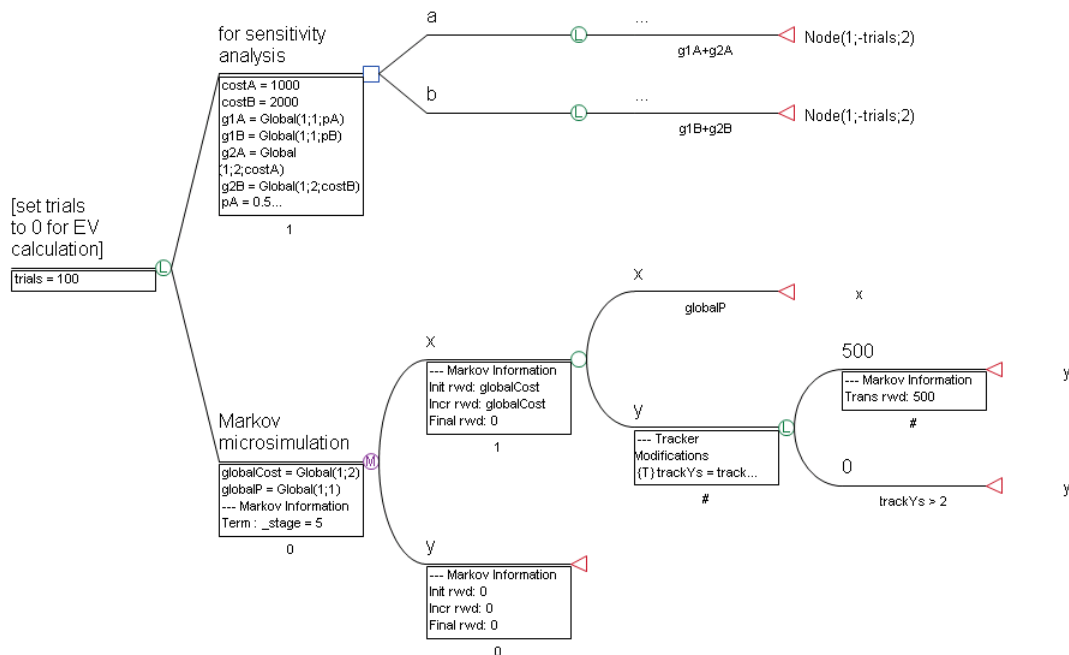
By placing a 1 in a Markov transition reward for an extra reward set, you can count the percentage of the cohort that passes through a specific transition. Be careful about your conclusions if the cohort could pass through that transition more than once.

35.10 Other advanced Markov options

35.10.1 Nesting or linking Markov models

TreeAge Pro includes powerful functions – for example, `Node()`, `Global()`, `GlobalIN()`, and `User()` — that can perform tasks such as: nest one Markov model within another, link multiple terminal nodes to one Markov model, or combine a Markov microsimulation with expected value analyses like 1-way sensitivity analysis. Function syntax is described in more detail in the Building Formulas Using Variables and Functions Chapter and the Tools and Functions for Complex Trees Chapter.

The Special Features tutorial example tree "Node Function" illustrates the use of these two functions.



Node Function Tree

The top arm is set up for an expected value calculation and its decision node includes two strategies both linked via the Node() function to the same Markov model at the bottom. Each strategy passes distinct values to the Markov process using the Global() function. The Node() function then calculates the Markov model using microsimulation trials (even during non-simulation analyses of the top decision node).

The `Node()` function's arguments are used to select a node in the tree and to determine what kind of calculation to use at that node. The syntax is:

Node(attribute; method; branch; ...)

In CE models, the *attribute* argument determines whether to return cost or effectiveness (-1 calculates and returns cost; -2 returns effect from previous calculation; -11 returns cost from previous calculation; -12 calculates and returns effect). In non-CE models, as in the example, use any number other than 0 (which returns 0).

For the *method* argument, use 0 to calculate the expected value; or specify a negative number to average that number of microsimulation trials (e.g., -100 to run 100 trials), as in the example.

The third and subsequent *branch* arguments are branch numbers used to select a node starting with a branch of the root node.

In the tree, the complete expression *Node(1;-trials;2)* returns the cumulative reward (attribute 1), based on the average of 100 trials (method -100) at the *Markov microsimulation* node (branch 2).

The `Global()` function is illustrated in this example as well. However, it is not a required element of linking Markov trees. Global matrices can be used to store transient values when it is helpful, such

as when you want to report the changing values of simulation tracker variables used in a subsidiary Markov process.

The following syntax is used to calculate and store a value in a cell in the Global matrix:

Global(row; column; expression)

The function returns the value of the calculated expression, as well as stores it in the Global matrix. The first cell in the global matrix is at row=1, column=1. Up to ten thousand cells are currently supported.

The following syntax can be used to reference a value saved to the global matrix:

Global(row; column)

The contents of the global matrix can be dynamically saved to a text file (or emptied) using the third syntax of the Global() function:

Global(value)

If the value evaluates to a non-zero number, the contents of the global matrix are silently saved to a text file in the tree's directory. A zero value will empty the matrix. Refer to the Tools and Functions for Complex Trees Chapter for more information.

In the example model, the variable definitions for *g1A*, *g1B*, *g2A* and *g2B* store information in the Global matrix. The definition *g1A = Global(1;1;pA)* stores the value *pA* in row 1, column 1 of the Global matrix.

At the unnamed branch of node *a*, the *g1a* and *g2a* are executed to generate a non-zero probability for the logic node, thereby placing the "*a*-related" values in the Global matrix. For the *b* strategy, the "*b*-related" values are placed in the Global matrix.

When the Markov model is evaluated (via Microsimulation) for each strategy by the Node function, the appropriate values are used when referenced in the Global matrix.

35.10.2 Additional notes on the Node() function:

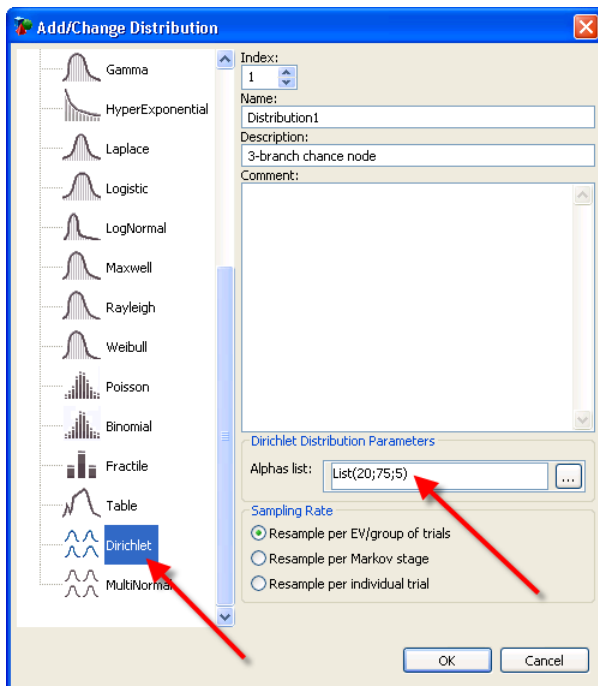
- Node() function syntax is described in more detail in the Tools and Functions for Complex Trees Chapter. The related Tree() and Global() functions are also covered in detail there.
- Trackers are not reset to 0 at the beginning of trials run by the Node() function. This is intentional in order to allow communication between the calling and called nodes.
- Trials run by the Node() function do not resample distributions automatically (even if their properties are set to sample per trial). To force a sample, use the Dist(N; 1) syntax or DistForce(N). To control the sampling rate, put the force sample expression in a tracker evaluation (outside a Markov process, for example).
- The second parameter of the Node() function, possibly specifying a number of simulation trials to be run, can be given a number after a decimal place to indicate that a statistical measure other than the mean should be used from the set of trials. Refer to the "Tools" Chapter.

35.10.3 Sampling probabilities from a multivariate Dirichlet distribution

If a chance node has more than two branches with non-negligible probabilities, performing a sensitivity analysis or Monte Carlo simulation that changes the values of these probabilities can be problematic. One option is to normalize the chance node's probability expressions. For example, if a node has three outcomes, A, B, and C, rather than assigning variables to two probabilities and using the # remainder for the third, you could do the following: assign three expressions that always sum to 1.0, such as $pA/(pA+pB+pC)$ and $pB/(pA+pB+pC)$ and $pC/(pA+pB+pC)$. No matter what values (≥ 0) are assigned to pA , pB , and pC , the three normalized probabilities will always sum to 1.0. (The # remainder could still be used in place of one of these.)

TreeAge Pro offers a similar solution using a special, multivariate form of the beta probability distribution, called a *Dirichlet distribution*. This distribution can be used to represent the uncertainty in all of the probabilities of a chance event. During Monte Carlo simulation, the distribution can sample probabilities for each branch using normalization to ensure that the probabilities always sum to 1.0.

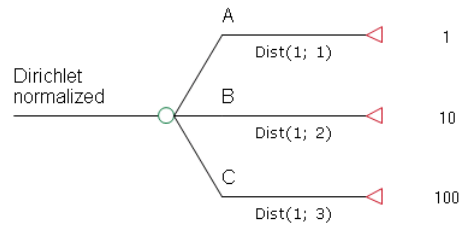
If the distribution is parameterized with a list of three positive *alpha* values, as shown here, TreeAge Pro will sample three independent $\text{Gamma}[\text{alpha}, \text{beta}=1.0]$ distributions and normalize these to create a list of three probabilities.



Dirichlet Distribution

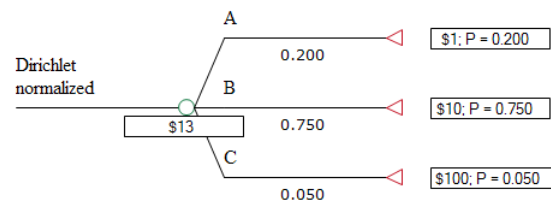
For documentation on creating and editing distributions, refer to the Monte Carlo Simulation, Distributions and Probabilistic Sensitivity Analysis Chapter.

The `Dist()` function is used with a second argument to indicate which branch (i.e., alpha) to use. The Health Care tutorial example model, "Dirichlet Simple", is shown below.



Dirichlet Simple Tree

Rolling back the tree shows the mean values of the probabilities, which are simply the normalized alpha parameters.



Dirichlet Simple Tree - rolled back

Performing a simulation in the example tree shows the effect of sampling independent Gamma distribution values based on the list of alpha parameters, and then normalizing. For each iteration of the simulation, a different set of Gamma random variates is drawn. Each iteration results in a different sum, as well as different ratios of the Gamma random variates to the sum (i.e., the probabilities), but normalization ensures that the resulting probabilities sum to 1.0.

The following output from a PSA simulation shows the different probability values generated by the Dirichlet distribution for each simulation iteration. Note that the probability values are centered around the respective mean values.

ITERATION	STRATEGY_1	DIST_1_VAR_1	DIST_1_VAR_2	DIST_1_VAR_3
1	11.478	0.197	0.767	0.036
2	11.946	0.158	0.805	0.037
3	13.086	0.190	0.757	0.053
4	15.122	0.225	0.696	0.079
5	11.276	0.225	0.738	0.037
6	12.008	0.131	0.834	0.035
7	13.733	0.188	0.752	0.060
8	11.489	0.258	0.700	0.042
9	14.635	0.193	0.736	0.071
10	12.317	0.168	0.789	0.043
11	12.178	0.165	0.794	0.041
12	12.439	0.235	0.715	0.051
13	11.734	0.142	0.824	0.033
14	10.197	0.279	0.691	0.030
15	13.635	0.163	0.780	0.057

PSA output - Values, Dists, Trackers

35.10.4 Markov cohort analysis using the TreeAge Pro Object Interface

Users of the TreeAge Pro Object Interface can create macros or other automation scripts/programs that run Markov analyses programmatically. For example, the macros would use a

TreeAgeProLib.ApplicationObj variable to create a TreeAgeProLib.TreeObj variable, and then a TreeAgeProLib.MarkovOutput variable.

The advantages of the programmatic approach to running the Markov cohort analysis include:

- Automate repetitive analyses
- Parse sections of the full text report for specific values
- Pause between cycles in order to make complex adjustments to the model (using MarkovEvents)

Refer to the Using the TreeAge Pro Object Interface Chapter for more information.

36. Individual-Level Simulation and Markov Models

This chapter covers the use of Monte Carlo simulation to run individual-level, discrete simulations (a.k.a. microsimulation) on Markov models. It covers topics including: tracker variables, 2-dimensional simulations (probabilistic sensitivity analysis), individual-level distribution sampling, parallel trials, dynamic simulation populations, and discrete event simulation.

Refer to the Analyzing Decision Trees Chapter and the Monte Carlo Simulation Chapter for general instructions on using Monte Carlo simulation. Refer to the Distribution Functions, Options and Types Chapter for details on using distributions for discrete simulation as well as probabilistic sensitivity analysis. The Cost-Effectiveness Simulation Reports and Graphs Chapter deals with the interpretation of Cost-Effectiveness simulation outputs.

36.1 Notes on simulation terminology

As described in the Monte Carlo Simulation Chapter, one important use of Monte Carlo tools in TreeAge Pro is in performing probabilistic sensitivity analysis (PSA). This chapter deals with a very different application of Monte Carlo analysis referred to variously as:

- Microsimulation – common name in this text
- Discrete (event) simulation
- 1st-order trials
- Individual-level simulation

In TreeAge Pro, these all refer to running the “Analysis > Monte Carlo Simulation > Trials (Microsimulation)...” menu command on a tree.

Microsimulation is often used with trees that include one or more Markov nodes. The function of simulation in such cases is to allow Markov models to evolve from the simpler cohort analysis approach to the more complex possibilities allowed by individual-level simulation. Plus, microsimulation can be used with *any* model; it is not restricted to trees including Markov nodes, and does not require the Healthcare module. For example, microsimulation can be used in combination with distributions that represent *variability* (i.e., use a continuous distribution for a rate of return in place of a chance node representing high, medium, and low returns).

This chapter focuses on many important aspects of building simulation models, including:

- Tracker variables (in Markov models)
- Distributions simulating individual-level variability
- Combining PSA and microsimulation
- Parallel trials and discrete event simulation

For more resources on simulation modeling, search this manual or use the Google® site search at:

<http://www.treeage.com/search.htm>

36.2 Simulation and tracker variables

As described in the previous two chapters, a Markov cohort (i.e., expected value) analysis retains no memory of previous events from one cycle to the next. Transitions and rewards are assigned to portions of the cohort based only on their state membership in the *current* cycle. The part of the cohort starting a cycle in a state is treated as homogenous, with no information about the different paths that lead to being in that state at that time.

However, to gain greater flexibility, modeling studies are increasingly using individual- or patient-level simulations (also referred to as discrete event simulation/DES or microsimulation) instead of standard Markov cohort analyses:

“The major drawback to Markov models is that they may not be suitable to tracking patients’ disease history properly, unless the analyst defines multiple health states, which may lead to intractable situations. They are also too rigid to take into consideration multiple patient-specific sociodemographic characteristics in a single model.... [Simulation resolves] these weaknesses and its flexibility allows patients with differing attributes to move from one event to another in sequential order while simultaneously taking into account important risk factors such as age, gender, disease history....”

Le Lay, “Can discrete event simulation be of use in modeling major depression?” *Cost Eff Resour Alloc.* 2006; 4: 19.

Some aspects of microsimulation in TreeAge:

- *Variables* can be used to store *state* (instead of adding additional state branches to the Markov node).
- *Simulation models* can avoid the restrictions of a fixed cycle length (i.e., the `_stage` counter) by defining a tracker variable to keep track of patient time.
- *Individual-level characteristics* – e.g., patient risk factors – can be sampled or bootstrapped from distributions or tables. This is covered later in this chapter.

36.2.1 Applications of tracker variables

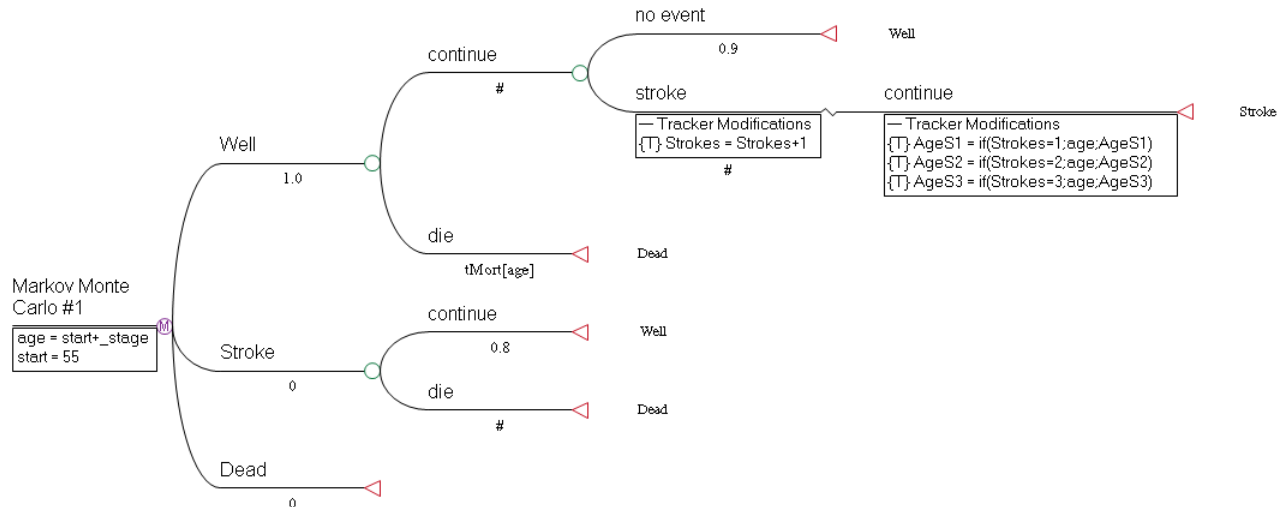
Tracker variables offer a simple way for the model builder to record any number of events while simulating individuals’ “runs” through the Markov model. For example, trackers can count the number of times an individual experiences important events, and even the timing of those events. A tracker variable’s “memory” of prior events can be easily updated, recalled, and reported. Tracker variables can also be used for recording and reporting extra “payoffs” (i.e., other than cost and effectiveness), without changing tree preferences, etc.

The value of a tracker variable can be updated simply by redefining the tracker variable at an event node, similar to the way that a regular probability or payoff variable/parameter is defined. For example, if a simulation trial encounters a node with the tracker modification:

$\{T\} \text{ Strokes} = \text{Strokes} + 1$

The current value of Strokes for that patient's trial is incremented by 1.

The Health Care tutorial example model “Markov Monte Carlo #1 - with table” is shown below.



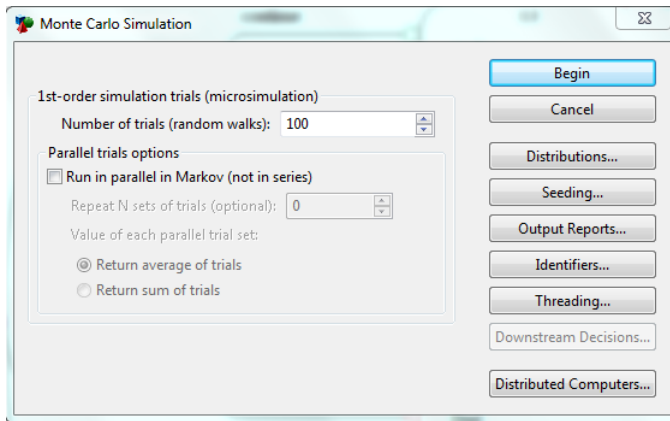
Markov Monte Carlo # 1 Tree

This model illustrates using trackers for recording individual history. The tracker variable *Strokes* counts the number of times an individual trial experiences the stroke event. Three other trackers record the age of the subject at the time of each successive stroke.

When running a microsimulation, as opposed to probabilistic sensitivity analysis, enter a number of trials, but leave the number of samples at zero.

To run a microsimulation:

- Select the root node.
- Choose Analysis > Monte Carlo Simulation > Trials (Microsimulation)... from the menu.
- Enter 100 trials in the Monte Carlo Simulation dialog.



Run Microsimulation

After the simulation is complete, you will be presented with summary statistics from the simulation output.

Statistic	Markov Monte Carlo #1
Mean	17.8
Std Deviation	13.4
Minimum	0.5
2.5%	1.5
10%	2.5
Median	15.5
90%	41.5
97.5%	46.0
Maximum	46.0
Sum	1,783.0
Size (n)	100.0
Variance	180.1
Variance/Size	1.8
SQRT[Variance/Size]	1.3

Actions
 ▶ Report Options...
 ▼ Text Reports
[Identifying Variables](#)
[Statistics - Values, Dists, Trackers](#)
 — Reports - Expanding —
[Strategy Values](#)
[Trackers](#)
 — Reports - Flat —
[Expected Values](#)
[Values, Dists, Trackers](#)
[Values, Dists, Trackers \(Tab-Delimited XLS/TXT\)](#)
[Trackers](#)
 ▶ Charts

Microsimulation output



Usually, the most critical statistic from a microsimulation is the *mean* value (one per strategy if run at a decision node). As described in the Building and Analyzing Markov Models Chapter, the mean value represents an *estimate of expected value* for the model and/or strategy. In many cases, the individual trial data is not valuable except as a contribution to the overall estimate of expected value. As you increase the number of trials in the microsimulation, you get better and better approximations of expected value.



Note that by default, several secondary output graphs are suppressed in Microsimulation output. Any output which relies on an iteration-by-iteration incremental comparison between strategies is not valid for Microsimulation. Such output is only valid when each iteration of the simulation is itself an expected value, as in the case of Probabilistic Sensitivity Analysis. If you need those outputs from Microsimulation (unlikely), it is possible to change the Tree Preferences (category Analysis Settings > Monte Carlo Options < Output Reports) by unchecking the box "Hide PSA reports for microsimulation."

At the start of each individual trial, all trackers are reset to 0. (This initial value can be changed in the variable Properties.) If an individual experiences the stroke event during the simulation, the tracker modification $Strokes = Strokes + 1$ is executed, incrementing the stroke counter. The *AgeS1*, *AgeS2* and *AgeS3* trackers then use conditional logic to store age at time of stroke 1, 2, and 3. The microsimulation's Values, Dists, Trackers link opens a text report which shows the final values of all trackers for each individual trail. (Each simulation will, by default, result in a different set of trials/individuals; refer to the Monte Carlo Simulation Chapter for more details.)

ITERATION	STRATEGY_1	AGES1	AGES2	AGES3	STROKES
1	15.5	60	64	69	3
2	6.5	0	0	0	0
3	10.5	59	0	0	1
4	16.5	56	67	0	2
5	7.5	61	0	0	1
6	46	69	75	99	3
7	3.5	57	0	0	1
8	27.5	59	63	0	2
9	23.5	61	66	77	3
10	1.5	0	0	0	0
11	14.5	57	64	68	3
12	10.5	0	0	0	0
13	8.5	62	0	0	1
14	37.5	62	67	70	5
15	3.5	57	0	0	1
16	13.5	55	67	0	2
17	46	99	0	0	1
18	23.5	61	65	0	2
19	9.5	56	0	0	1

Microsimulation output - Values, Dists, Trackers

36.2.2 Referencing trackers in transition probabilities and other Markov calculations

The example shown in the previous pages uses trackers for reporting only – i.e., {T} Strokes does not affect the Markov calculations. Sensitivity analysis, Markov cohort analysis, and other expected value calculations can still be used with this model – *trackers will simply be ignored*.

In addition to their uses in reporting, trackers have important applications as parameters in calculations of transition probabilities, rewards, 1st-order distributions, termination conditions, or any other tree expressions. In other words, the trackers are not limited to recording events; they can also be used to affect future transitions, events, rewards, etc.

For example, one set of tracker variables could be used in a model to indicate how often a patient has undergone a particular treatment for cancer, while another set of tracker variables keeps track of the current size, type, and location of a tumor. A logic node could compare the value of the tumor size tracker variable to some threshold and transition the individual to the symptomatic state. Or, the tumor location tracker variable could be used as a lookup value in retrieving an appropriate Markov reward from a table of surgical costs.

In the Markov Monte Carlo # 1 Tree shown above, perhaps the probability of death is affected more by the number of strokes than by age. In that case, you could set up a new table and reference the *Strokes* tracker in the probability of death with a new table expression like this one.

Probability: *TblDeathStrokes[Strokes]*

Once model *calculations require trackers* in order to calculate correctly, the model should be analyzed *only with microsimulation trials*. Expected value analyses (i.e., roll back, n-way sensitivity analysis, Markov cohort analysis) will ignore the trackers and will probably yield incorrect results.

Regular n-way sensitivity analysis is not generally used with models requiring microsimulation/trackers. Instead, a Monte Carlo probabilistic sensitivity analysis (PSA) with an inner microsimulation (a 2-dimensional simulation) might be used to examine parameter uncertainty.

36.2.3 Creating and defining tracker variables

Tracker variables are created and defined in a similar way as the regular variables described in the variables chapter; however, you create and edit trackers in the Tracker Properties and Tracker Modifications Views rather than their equivalent variable views.

Trackers also differ from variables in the way they are evaluated during analyses. This will be discussed further within this chapter.

One quick way to create and define a tracker is by right-clicking on a node that requires a tracker modification.

To create and define a Monte Carlo tracker variable:

- Right-click on an event node, usually within the Markov subtree, where you want the tracker modification.
- Choose Define Tracker > New Tracker from the context menu.
- In the Add/Change Tracker dialog, enter a name and other properties of the tracker and click OK to save it.
- In the Define Tracker dialog, enter the tracker modification and click OK to save it.

The screenshot shows a Windows-style dialog box titled "Add/Change Tracker". It has two tabs: "General" (selected) and "Categories". The "General" tab contains the following fields and controls:

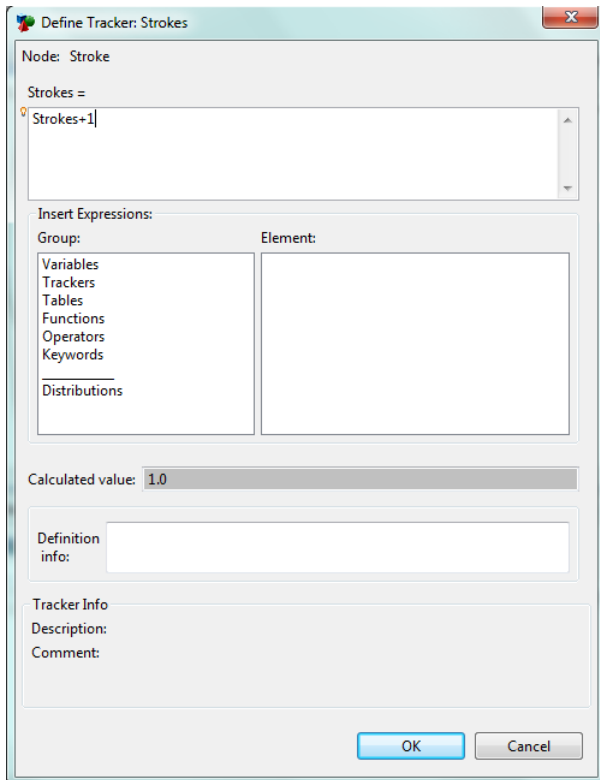
- Name:** A text box containing "Strokes".
- Description:** A text box containing "Number of strokes experienced by trial".
- Comment:** A large, empty text area.
- Show modifications in tree:** A checked checkbox.
- Initial value:** A text box containing "0".
- Evaluation priority: (0=first):** A spin box containing "0".

At the bottom right of the dialog are "OK" and "Cancel" buttons.

Add/Change Tracker Dialog

The initial value is the value of that tracker variable at the beginning of each trial. In most cases, the default value of 0 is appropriate, while in rare cases, a different value is more appropriate. For example, let's say you want to record the `_stage` value when a certain transition occurs. If the initial value is zero, there will be no way to distinguish between the case where the transition occurred at `_stage` 0 and the case where the transition never occurred. An initial value of -1 would distinguish between those cases.

The evaluation priority value determines the order in which to evaluate trackers at a node. This only impacts analysis results if you have a tracker that is updated and used in another tracker modification at the same node. In such a case, you might want to ensure that the updated tracker value is used in the other tracker modification. The priority can be set for each tracker; however, the priority is only used if the appropriate Tree Preferences option is set in the Other Calc Settings category.



Define Tracker Dialog

A tracker modification like this...

{T} Strokes=Strokes+1

... is typical for counting an event. Note that the text you actually enter in the Define Variable window is only what appears to the right of the equals sign in the example above.

When TreeAge Pro displays a tracker modification in the tree view or Define Variable window, it appears with the prefix "{T}". Note that you should not type the {T} prefix when referring to the tracker in formulas.

To change (or delete) an existing tracker modification, simply right-click on the desired event node and choose the variable from the Define Variable list. Or, use the tracker views described in the next two sections.

36.3 Tracker views and dialogs

This section describes the views and dialogs related to trackers.

36.3.1 Tracker Properties View

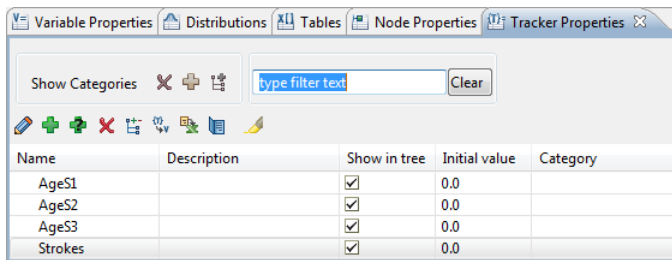
The Tracker Properties View is used to manage tracker properties. Unlike tracker modifications, tracker properties apply to the entire tree. Therefore, the Tracker Properties View is a tree-level view rather

than a node-level view. This view functions in a manner similar to the Variable Properties View, except it is used for trackers.

To open the Tracker Properties View:

- Choose Views > Tracker Properties from the toolbar.

Below is an image of the Tracker Properties View from the Health Care tutorial example model "Markov Monte Carlo #1".



Tracker Properties View

The main grid contains a list of the tree's trackers along with a few of the key variable properties. These properties can be edited within the grid. If you change the tracker name, all references to that tracker within the tree will be modified as well.

Filter and category options work the same as for variables. Refer to the section describing the Variable Properties View for details.

The Tracker Properties View toolbar provides additional functions.



Tracker Properties View toolbar

The functions associated with the icons are presented below from left to right/top to bottom.

1. Edit tracker
2. Add new tracker
3. Add multiple new trackers
4. Delete selected tracker
5. Group trackers by categories
6. Convert tracker to variable
7. Edit in Excel
8. Report
9. Highlight

Tracker Properties View toolbar functions

These functions are described briefly here. Since the functions are nearly identical to the functions for editing regular variables, please refer to the Working With Variables Chapter for more details.

Edit tracker: Edit the properties of an existing tracker via the Add/Change Tracker Dialog.

Add new tracker: Create a new tracker and enter its properties via the Add/Change Tracker Dialog.

Add new trackers: Specify the number of new trackers and then enter their properties via the Add/Change Tracker Dialog.

Delete selected tracker: Delete the tracker(s) selected in the view's grid.

Group trackers by categories: Group the trackers by category in the view's grid.

Convert tracker to variable: Convert the selected tracker to a regular variable.

Edit in Excel: Use the Excel Module to edit tracker properties in Excel.

Report: Generate a report on trackers in the model.

Highlight: Highlight the tracker within the model in the Tree Diagram Editor.

36.3.2 Tracker Modifications View

The Tracker Modifications View is used to manage tracker modifications at different nodes within the tree. Tracker modifications are created at specific nodes, meaning the Tracker Modifications View is a node-level view. The contents of the view reflect the context of the selected node. This view functions in a manner similar to the Variable Definitions View, except it is used for trackers.

To open the Tracker Modifications View:

- Choose Views > Tracker Modifications from the toolbar.

Below is an image of the Tracker Modifications View from the Health Care tutorial example model "Markov Monte Carlo #1" with the *continue* node selected.

The screenshot shows the Tracker Modifications View for the "Markov Monte Carlo #1" model. The view displays a table with the following data:

Name	Definition	Info/Comment
---Defined---		
AgeS1	if(Strokes=1;age;AgeS1)	
AgeS2	if(Strokes=2;age;AgeS2)	
AgeS3	if(Strokes=3;age;AgeS3)	
---Undefined---		
Strokes		

Tracker Modifications View

Note that the three tracker modifications from the *continue* node are presented in the view, while the tracker *Strokes* is presented within the *--Undefined--* group.

The functions within this view work almost identically to the Variable Definitions View. Please refer to that section for more details. Note that tracker modifications cannot be inherited; that functionality from the Variable Definitions View is not included in the Tracker Modifications View.

36.3.3 Add/Edit Tracker Modification Dialog

The Add/Edit Tracker Modification Dialog is used to edit tracker modifications at a specific node. The dialog is described in an earlier section of this chapter.

36.4 More notes on trackers

36.4.1 Critical differences between trackers and variables

Although much of the content of this section has already been presented, it's very important to understand the differences between trackers and variables. Therefore, those differences are restated here:

Item	Trackers	Variables
Value scope	Applies to an individual trial.	Applies to the entire cohort.
Analysis type	Requires microsimulation.	No restrictions.
When are definitions/modifications executed?	Immediately at the node where the tracker modification is found.	Only when the variable is referenced in a required calculation (i.e., payoff, reward, probability, etc.).
Initial value	Required at the starting point for each trial.	Does not exist.

Differences between trackers and variables

36.4.2 Tracker modification location and timing

- A node's tracker modifications occur after that node is selected in a random walk (i.e., after any logic or probabilities for the branch and its siblings have been evaluated), and after rewards (state or transition) are accumulated at that node.
- Modifications will typically be defined at, or to the right of, a Markov node, as in the examples in this chapter. However, if any chance nodes precede a Markov node, tracker modifications can also be placed at these earlier events.
- Modifications at a Markov node will be evaluated once at the start of *_stage 0*, when a trial starts the Markov process.

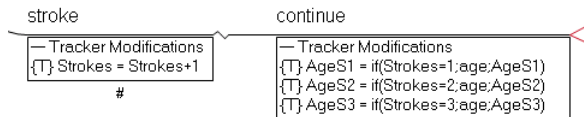
- Modifications at an absorbing state are only evaluated at initialization (at `_stage 0`), and are ignored later. Modifications at the root node of the tree will be ignored, unless it is a Markov node.

Notes on tracker modification location and timing

36.4.3 Tracker modifications that reference other trackers

If you have more than one tracker modification at a particular node, the modifications will be applied in *reverse alphabetical order*. To avoid errors and confusion, if `{T} TrackerB` is dependent upon the value of `{T} TrackerA`, it is recommended that they be defined at successive nodes. Simply move the dependent tracker modification to a node to the right.

This is accomplished by inserting a single branch to the right of the existing event, and moving the dependent tracker modification to that node. The node between the dependent modifications can be changed to a label node, as shown below.



Tracker Strokes set before it is used

36.4.4 Trackers and expected value calculations

While expected value (EV) analysis is not recommended for trees in which tracker variables are referenced in probability or reward calculations, this type of analysis is not disabled. Since tracker modifications are only meaningful within simulation trials, TreeAge Pro generally ignores them during EV calculations, *including Monte Carlo PSA*. Outside of simulation trials, the values of trackers will be equal to their initialization value.

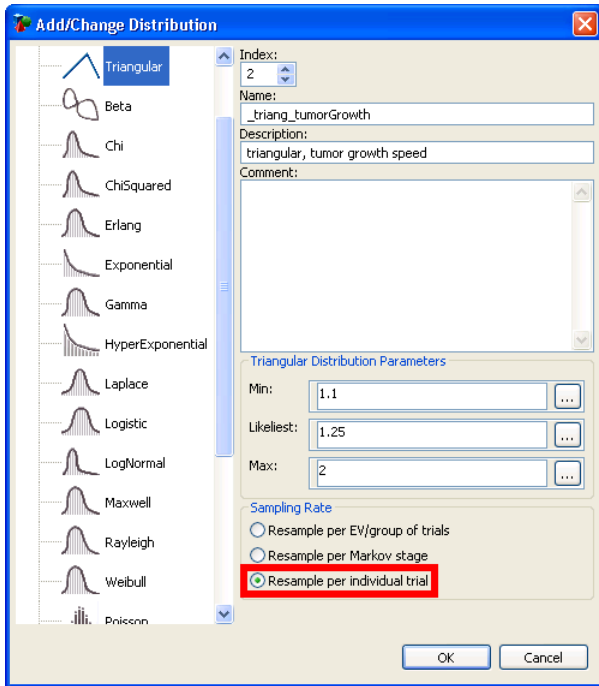
One exception to this rule is when an expected value analysis makes a call to the `Node()` function which in turn runs a set of microsimulation trials at another location in the tree. See the information on the `Node()` function in the Tools and Functions for Complex Trees Chapter and the Markov Modeling Tools and Techniques Chapter.

A more efficient replacement for recursive variable definitions is another possible application of trackers in expected value calculations of regular trees. In Tree Preferences, under Other Calc Settings, there is a preference that must be turned on for tracker modifications to work like recursive variable definitions during regular, non-microsimulation analyses.

In light of the potential for error, it is advisable to limit analysis of trees that require tracker variables to microsimulation only. For the purposes of EV analyses like roll back or one-way sensitivity analysis, it may be desirable to either use the advanced `Node()` linking function or to develop a modified version of the model in which calculations do not depend on tracker variables.

36.5 Sampling individual-level distributions during simulation

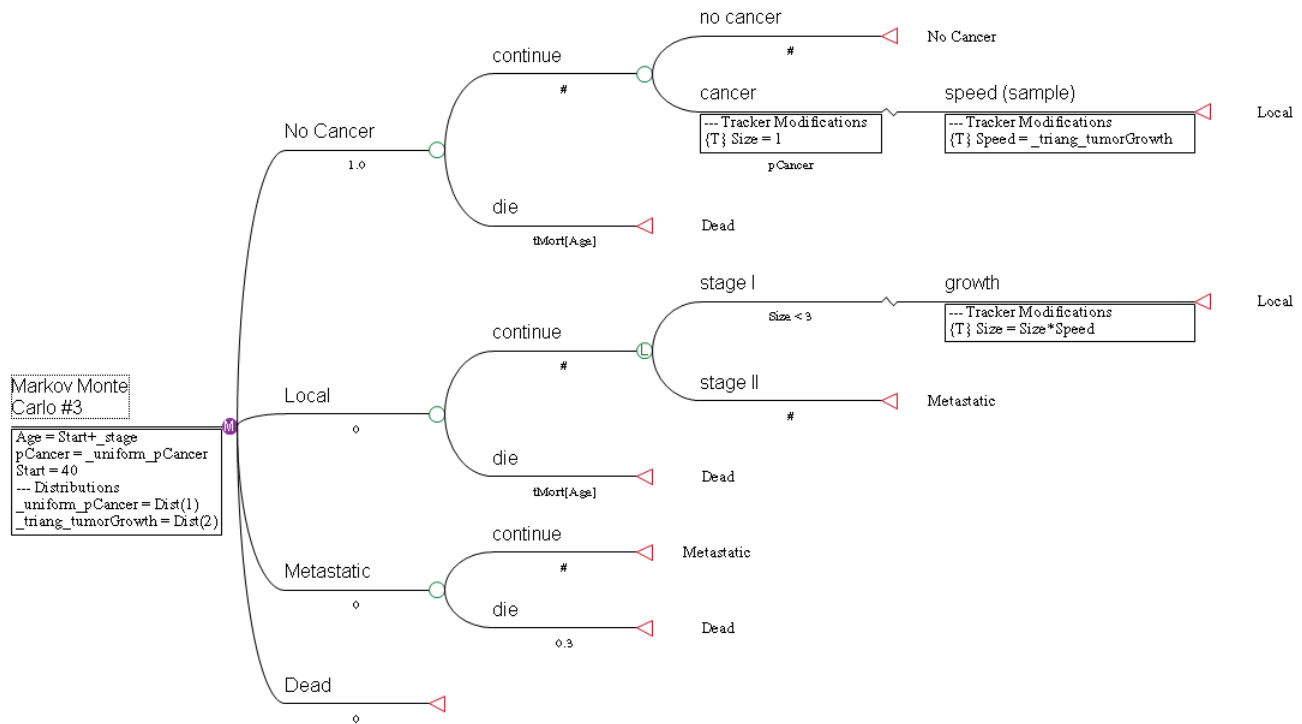
Rather than representing *parameter uncertainties* for probabilistic sensitivity analysis (PSA), some modeled distributions are used to represent *individual variability* in the context of microsimulation trials. Such distributions might be used to sample values that vary between individuals or from cycle to cycle. In TreeAge Pro, this is implemented by changing the sampling rate in the distribution's properties.



Sample by trial

During any microsimulation (including an inner loop within a 2-dimensional PSA), distributions that are changed from the default setting to "Resample per individual trial" will automatically sample at the appropriate rate during the simulation. Each individual (i.e., trial) will effectively get their own random sample.

The Health Care tutorial example model "Markov Monte Carlo #3" is shown below. In addition to using tracker variables, it also includes an individual-level distribution (#2) as well as a regular PSA-type distribution (#1). Microsimulation must be used to analyze the model; without it, tracker variable modifications are ignored and no cancer will ever metastasize in the model. Microsimulation-level sampling will also factor in variability of tumor growth rates.



Markov Monte Carlo #3 Tree

36.5.1 Notes on sampling 1st-order distributions

- Refer to the Monte Carlo Simulation, Distributions and Probabilistic Sensitivity Analysis Chapter for information on creating and managing distributions.
- To perform a probabilistic sensitivity analysis on a model, including both first-order and second-order distributions (like "Markov Monte Carlo #3"), the analysis should use 2-dimensional simulation.
- Refer to the Distribution Functions, Options and Types Chapter for information on sampling from tables. This approach can be used to "bootstrap" patients into the model from a table containing patient data/characteristics (e.g., one row per patient). The built-in keyword `_trial` can be used as a patient counter to pick the appropriate row predictably, or a uniform (integer-only) distribution can be used to separately sample a patient number for each trial. (In a 2-dimensional simulation, the `_sample` counter increments in the outer loop.)

36.6 Filtering/refactoring simulation output

TreeAge Pro allows for Monte Carlo simulation output to be filtered and/or refactored in the following ways.

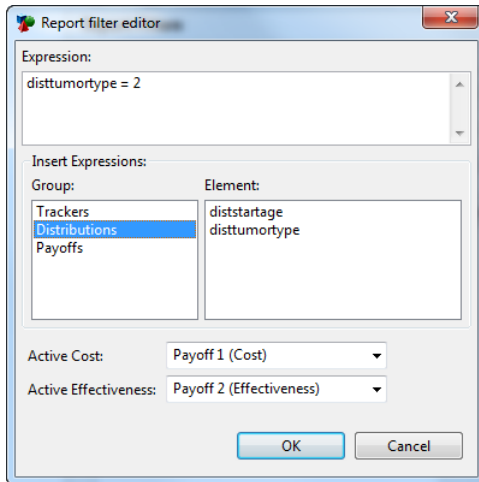
- Filter the output to include only iterations that meet selection criteria.
- Change the active payoffs.

A Preferences dialog opens to show any existing filters (of which there are none). Click Add to create a filter for editing in the Report filter editor, which provides you with the option to filter results based on any input characteristic (distributions) or output data (payoffs, trackers).



When filters use output data, strategies will not have the same results for each simulation iterations. If you apply a filter to output data, the row will remain in the filtered data set, if the iteration row meets the filter criteria for any of the strategies.

To filter for the more aggressive tumor type, the following filter would be applied.



Report filter editor

Note that in addition to providing access to the input and output data fields for filtering, the Report filter editor allows you to change the active payoffs for the model's calculation method. If you had multiple valuations of cost and/or effectiveness, you could change the active payoffs here to reevaluate the model (CE analysis) using different model outcomes.

Once the filter is complete, click OK. Then select the filter in the Preferences dialog, choose Set Active, and click Apply.

Attribute	Statistic	Existing Tx	New Tx
Cost	Mean	307,589.49	418,883.42
	Std Deviation	201,688.45	270,288.42
	Minimum	10,000.00	10,000.00
	2.5%	30,000.00	30,000.00
	10%	70,000.00	90,000.00
	Median	270,000.00	370,000.00
	90%	600,000.00	810,000.00
	97.5%	780,000.00	1,030,000.00
	Maximum	985,000.00	1,180,000.00
	Sum (n*Mean)	0.00	0.00
	Size (n)	3,045.00	3,045.00
	Variance	40,678,228,855.08	73,055,830,414.61
	Variance/Size	13,359,024.25	23,992,062.53
	SQRT[Variance/Size]	3,655.00	4,898.17
Eff	Mean	7.87	8.86
	Std Deviation	4.75	5.04
	Minimum	0.48	0.48
	2.5%	1.38	1.38
	10%	2.28	2.38
	Median	6.98	8.08
	90%	15.08	16.88
	97.5%	18.52	18.57
	Maximum	19.00	19.00
	Sum (n*Mean)	0.00	0.00
	Size (n)	3,045.00	3,045.00
	Variance	22.52	25.41
	Variance/Size	0.01	0.01
	SQRT[Variance/Size]	0.09	0.09

Microsimulation results filtered

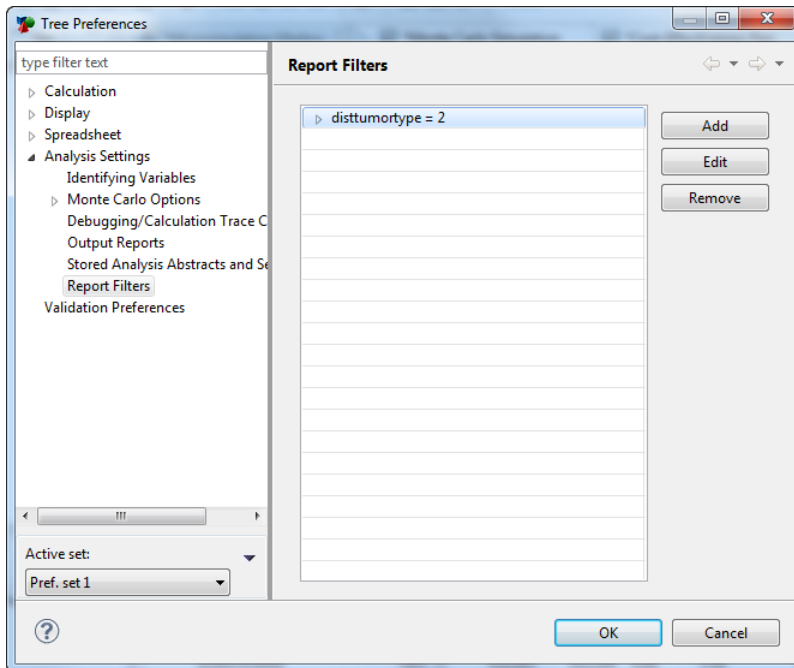
Note that the total number of iterations has been reduced from 10,000 to 3,045, reflecting only the iterations with tumor type 2. All secondary outputs, including the CE Rankings report below will now be generated from the filtered results. See below.

Category	Strategy	Cost	Incr Cost	Eff	Incr eff	Incr C/E (ICER)
Excluding dominated	Existing Tx	307,589		7.87		
	New Tx	418,883	111,294	8.86	0.98	113,093

Microsimulation results filtered - CE Rankings

Note that in the filtered results for the more aggressive tumor, the overall effectiveness for each strategy is about 1.4 QALYs less. However, the ICER does not change much, so the strategy selections would likely be the same. This may not always be the case.

When a filter is created for simulation output, the new filter is automatically copied back into the Tree Preferences for the model, so it can be reused for future analyses. See below.



Filter stored in tree preferences

36.7 Debugging simulations, the Calculation Trace Console

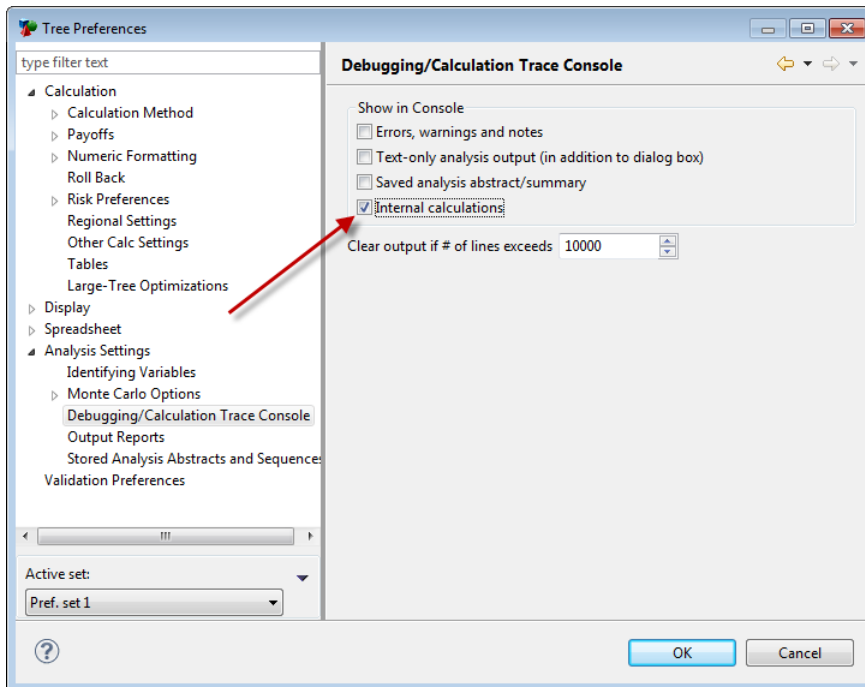
The Calculation Trace Console can be used to display/report a variety of textual information to help debug a model, including: error messages, text-only analysis output, stored analysis summaries, and detailed internal calculations (e.g., step-by-step evaluation of variables).



In previous versions of TreeAge Pro, debug output was written to the Debug Pane. This output is now written to the Calculation Trace Console.

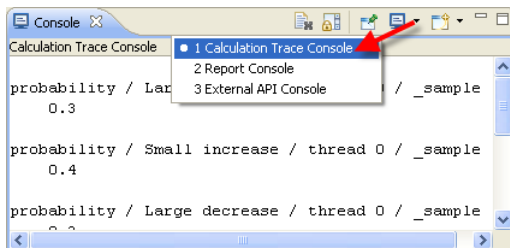
To turn on the output of debug information:

- Choose Tree > Tree Preferences from the menu or press *F11*.
- Navigate to the Tree Preferences category Analysis Settings > Debugging/Calculation Trace Console.
- Check boxes to send certain output to the Calculation Trace Console.



Tree Preferences - Debugging

The fourth option, “Internal calculations” is potentially of great use when debugging simulations and other complex analyses and models. If the “Internal calculations” debugging preference is turned on in a tree, each variable, probability, and other calculation is reported in detail in the Calculation Trace Console View.



Calculation Trace Console View

During simulations, random walks and distribution samples are also detailed. The Calculation Trace Console View output can then be used to check calculations performed during the analysis.

The figure below shows some of the calculation output from a Microsimulation run on the Markov Monte Carlo #3 model.

- Falls in the range for branch *continue*
 - Walk to *continue*
4. Repeat this process for the next random walk to *no cancer* or *cancer*.

Calculation trace output

36.7.1 Using the Debug() function

The Debug() function can be used to control/limit debugging output during simulations, or to add custom text to the pane. This may be of interest when an internal calculation debugging output may be millions of lines per analysis. Use the Debug() function to dynamically turn on and off the flow of calculation outputs to the Calculation Trace Console at strategic points in the analysis.

Refer to the Tools and Functions for Complex Trees Chapter for details on the Debug() function syntax.

36.7.2 Using the GlobalN() function

Another more customizable approach to debugging complex simulations involves the Global matrices. The GlobalN() and Global() functions can be used in a variety of ways, and are capable of storing millions of user-specified values and calculation results during simulations and other analyses.

The section on parallel trials and dynamic microsimulation later in this chapter provide additional discussion about using the GlobalN function for reporting. For more details, refer to the Tools and Functions for Complex Trees Chapter.

36.8 Parallel trials, discrete event simulation, and dynamic populations

Traditionally, microsimulation trials in TreeAge Pro are run in *series*. In other words, when trial #1 finishes (e.g., a Markov process terminates), then trial #2 begins. Added in TreeAge Pro 2004, there is an option that exists to run Markov microsimulation trials in *parallel* instead. In parallel trials, all *n* trials are stepped through a Markov process one cycle or event at a time, rather than running each trial to completion before starting another.

Parallel trials were introduced initially to enable the StateProb() function to work during microsimulation in much the same way it does during a cohort analysis – returning the percentage of the model “population” that started the current cycle in a particular state. Refer to the section on the StateProb() function from the previous chapter.

Recent versions of TreeAge Pro have added more features that extend the capabilities of parallel trials. For example, GlobalN() and related functions create “public” space for communicating data, such as a disease state, between different simulation trials (versus trackers, which function as private information within a single trial).

The following sections cover the basic functionality of parallel trials simulations, including some example references. It also describes new parallel trials features that were added in TreeAge Pro 2008:

- *Synchronized parallel trials* - triggered by a user-defined tracker variable, {T} _CLOCK, representing patient time.
- *Open/dynamic populations* - using non-coherent initial and transition probabilities to dynamically create individuals/trials.
- *Multiple sets of parallel trials* can be run, for example, to average out variability associated with a small/dynamic population.

A review of some simulation terms is also provided; the basic underlying message being that all microsimulations are discrete simulations. However, parallel trials simulation will be necessary to handle some aspects of discrete event simulation, including interaction and resource competition between “entities” (i.e., trials).

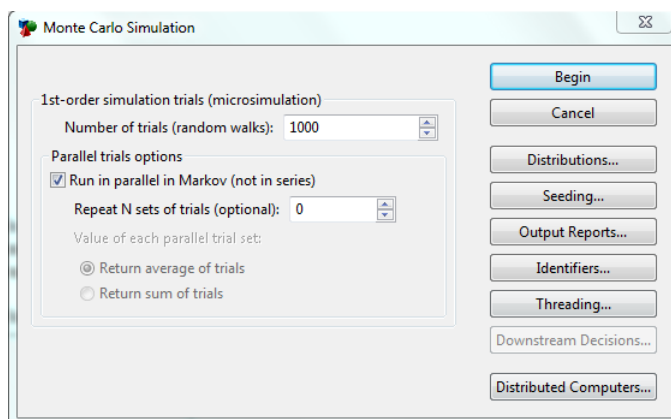
36.8.1 Running a parallel trials simulation

In many cases, no special changes are required to run parallel microsimulation trials on a model. Although, a certain basic tree structure is required (so some existing trees will not allow parallel trials).

Parallel trials are currently only available when running a simulation at either:

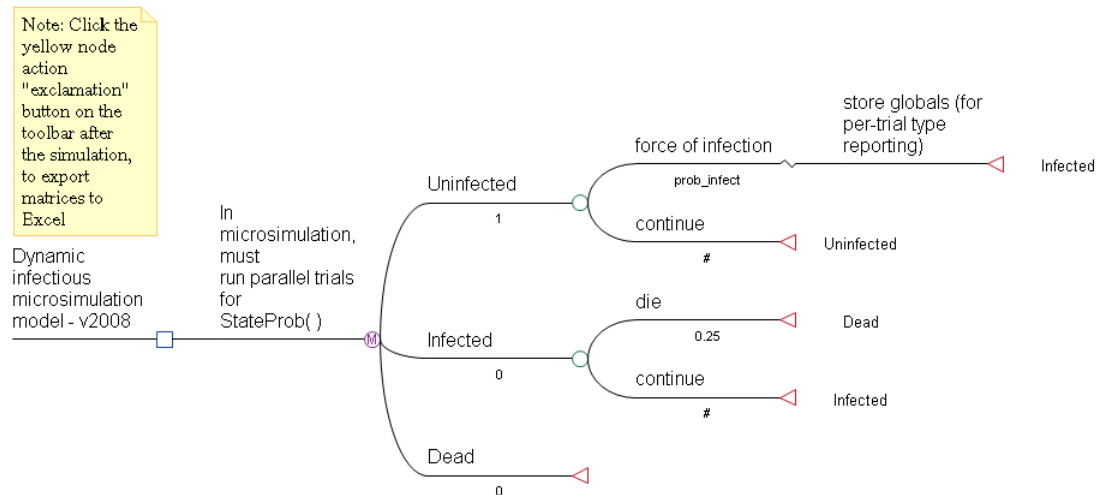
- a Markov node
- a decision node with only Markov branches
- a label node to the left of one of the above nodes

With an appropriate node selected, the microsimulation setup options include a “Run in parallel” checkbox below where you enter the number of trials to run. This option is also available in 2-dimensional simulations.



Setup microsimulation with parallel trials

The Health Care tutorial example model “Parallel Trials Closed Population” is illustrated below.



Parallel Trials Closed Population tree

The tree uses the StateProb() function to model a simplistic force of infection calculated based on the size of the infected versus total population. Note the following three variable definitions:

$sp_un = stateprob(1)$ - Get the percentage of the trials that are uninfected

$sp_inf = stateprob(2)$ - Get the percentage of the trials that are infected

$prob_infect = If(sp_un; .5 * sp_inf / (sp_un + sp_inf); 0.05)$ - Calculate the probability of infection based on the prior two percentages

The model can be calculated using cohort analysis (i.e., transitions to infected work correctly without trackers). However, trackers are included to enable extra reporting if simulation is used; *for StateProb() to work, the simulation must use parallel trials.*

36.8.2 Discrete event simulation (DES)

Most Markov models have a fixed cycle length, creating a linear relationship between cycles and time. However, some modelers might find this relationship artificial as events in reality can happen at any point within the continuous passage of time.

When Markov models are analyzed via microsimulation, you can track time on a continuous basis by sampling the amount of time between events from distributions. This type of model is a Discrete Event Simulation (DES) model.

In general usage, discrete event simulations often model a few things not typical in microsimulation:

- Entity (e.g., patient) time and/or system time modeled as stochastic jumps (rather than fixed increments)
- Parallel processes (interacting and competing)
- Dynamic creation of entities



TreeAge Pro 2014 added new functionality with the new DES and Time node types. We recommend using this new functionality rather than tracking time in DES models with trackers. Refer to the Time-to-Event Chapter for details.

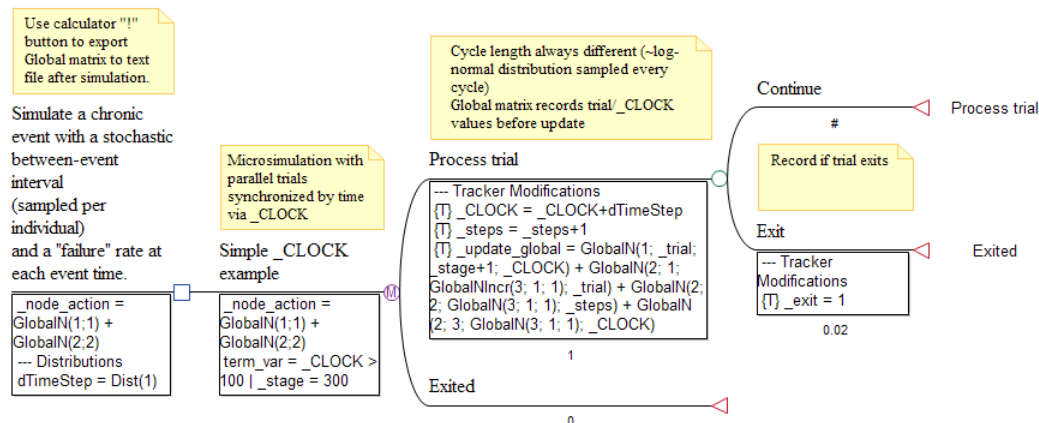
As described in the following sections, all of these can be handled in a microsimulation using parallel trials. (More complex Health Care tutorial example models – “Complex Parallel Trials v2008,” “Queue Problem”, et al – illustrate the relevant parallel trials features that will be shown.)

36.8.3 Synchronizing parallel trials using {T} _CLOCK

If your model defines a tracker called "{T} _CLOCK" in order to represent patient time and parallel trials are run, _CLOCK will be used to effectively “synchronize” the parallel trials.

Microsimulation trials (parallel or not) normally proceed under the standard Markov assumption of fixed cycle length. In the special case of a parallel trials simulation, individual trials progress one cycle at a time in an orderly fashion, with trial #1 completing a cycle, then trial #2, and so on. However, discrete event simulation models typically discard the _stage counter (along with fixed cycle length) in favor of using a tracker variable to store patient time.

In a non-parallel discrete simulation (in which trials run independently/in series), the Markov model can use a tracker variable with any name in order to store patient time. To store patient time in a tracker in a parallel trials simulation, however, a tracker called _CLOCK should be used. This is illustrated in the Special Features tutorial example tree “Parallel Trials _CLOCK #1”, shown below.



"Parallel Trials _CLOCK #1" tree

The example models different cycle lengths for different individuals (e.g., based on different risk factors), using a tracker to keep patient time instead of the _stage counter. Under the assumption that parallel trials were eventually going to be required – e.g., the StateProb function was going to be used – the special name _CLOCK is used for the patient-time tracker:

$\{T\} _CLOCK = _CLOCK + dTimeStep$

Since {T} _CLOCK exists, parallel trials will not proceed on a trial by trial basis. For example, after each trial has completed its initial cycle (i.e., _stage 0), trial #1 is *not* automatically run for a cycle. Instead, the trial with the lowest value of _CLOCK is run for one cycle. After this, another trial is selected in the same way, and so on. This process continues until all trials terminate (or a maximum number of total cycles are exceeded).

Look at the Values, Dists, Trackers output below that was generated from the parallel trials simulation.

Monte Carlo Simulation Report							
ITERATION	STRATEGY_1	_CLOCK	_EXIT	_STEPS	_UPDATE_GLOBAL	DIST_1	
1	0	105.9844132565	0	15	212.8375714121	7.0656275504	
2	0	56.9308756992	1	22	131.6862172439	2.5877670772	
3	0	111.343475148	0	8	204.851081509	13.9179343935	
4	0	49.6234115374	1	9	100.2193982887	5.513712393	
5	0	76.9166669148	1	10	152.4500004467	7.6916666915	
6	0	113.2572514757	0	7	206.1552882441	16.1796073537	
7	0	112.3321667031	0	9	214.7016296944	12.4813518559	
8	0	18.5228580793	1	5	41.6365729268	3.7045716159	
9	0	104.6641278609	0	18	223.6989081816	5.81467377	
10	0	68.3812872775	1	4	115.5719309162	17.0953218194	

Simulation output

Note that each trial that did not "Exit" (_exit = 1) has its own number of cycles (recorded in _steps); however, each stopped processing after the _CLOCK value exceeded 100.

The model also generates text output via the Global matrices 1 and 2. Global matrix 2 below shows the order in which trials were processed.

parallel trials_clock1_globaln_2																					
	A	B	C	D	E	F	G	H	I	J	K	L	M	N	O	P	Q	R	S	T	U
1	globaln(2)																				
2	C:\Eclipse\ eclipse351-treeage\branch11\src\ eclipse\plugins\com.treeage.treeagepro.tutorials\tutorials\Healthcare\Parallel Trials _CLOCK 1.trex																				
3																					
4	C1	C2	C3	C4	C5	C6	C7	C8	C9	C10	C11	C12	C13	C14	C15	C16	C17	C18	C19	C20	C21
5	1	2	3	4	5	6	7	8	9	10		2	8	2	4	9	1	8	5	2	4
6	0	0	0	0	0	0	0	0	0	0		1	1	2	1	1	1	2	1	3	2
7	0	0	0	0	0	0	0	0	0	0	0	2.587767	3.704572	5.175534	5.513712	5.814674	7.065628	7.409143	7.691667	7.763301	10.35107

Simulation output - Global matrix 2

Note that after each of the 10 trials run in order to start; the trial (row 5) with the smallest value of _CLOCK (row 7) is executed next, regardless of the number of steps/cycles (row 6).



The Evaluator View's "!" button executes the _node_action variable definition from the root node. By clicking this button immediately after the simulation (with the tree active in the editor), global matrices 1 and 2 are output to text files.

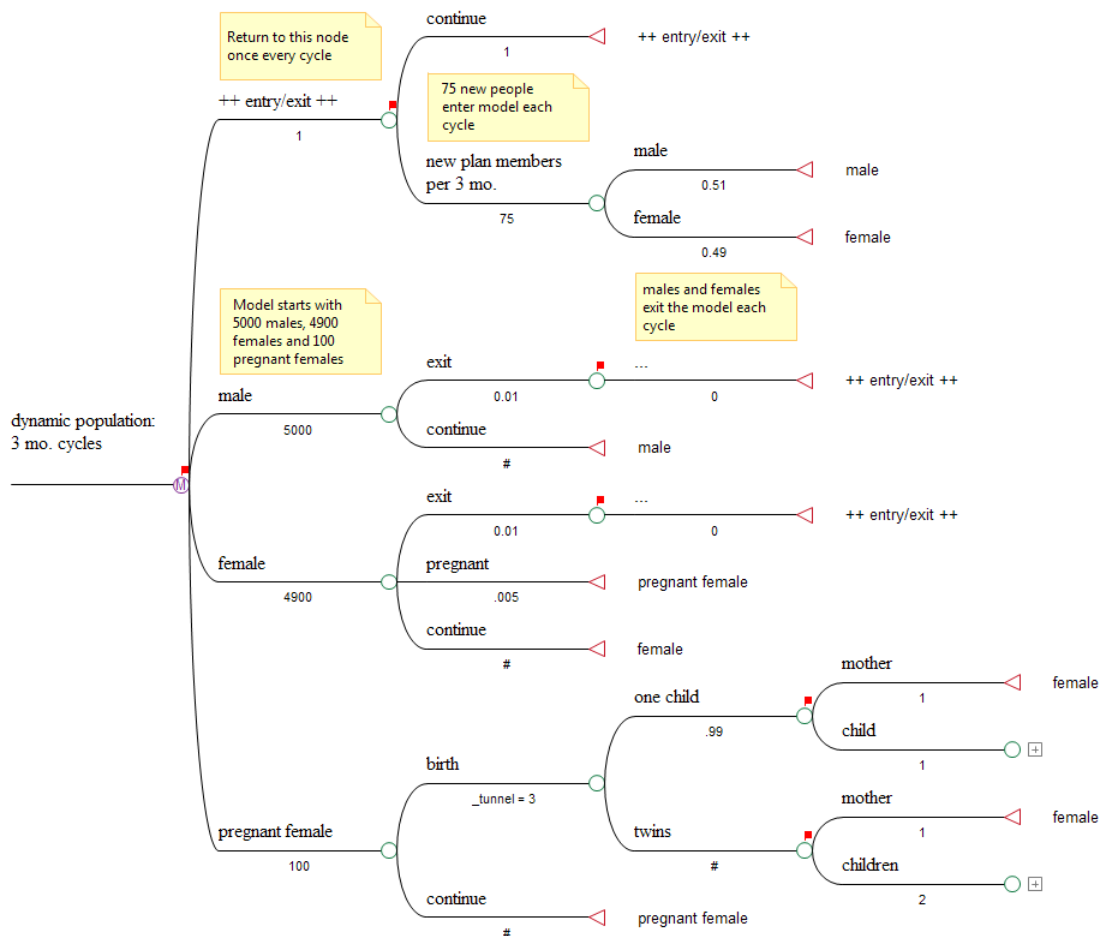
Refer to the remainder of this section for additional features related to {T} _CLOCK synchronization of parallel trials.

36.8.4 Parallel trials and open/dynamic populations

Prior to TreeAge Pro 2008, simulations required the regular pre-specified/fixed number of trials, as well as coherent probabilities. Now, however, dynamic/open populations can be simulated using non-coherent Markov initial and/or transition probabilities and parallel trials.

As described in the previous chapter, simulating open/dynamic populations with parallel trials utilizes the same basic approach as dynamic cohort analysis models. Simulation models can utilize non-coherent initial probabilities (to start a certain “number” of individuals in a particular state) and non-coherent transition probabilities (to “inject” an arbitrary number of additional members into any state at any time), provided that parallel trials are run.

The Health Care tutorial example healthcare tree “Dynamic Population v2008” used to illustrate dynamic cohort models in the previous chapter is shown again here.

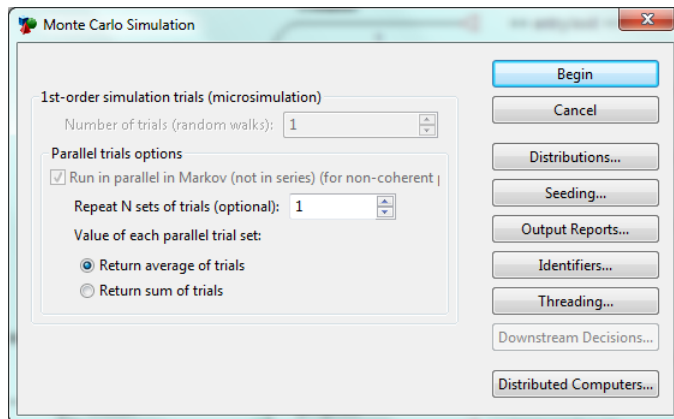


Dynamic population model

To illustrate the use of non-coherent Markov probabilities to model a dynamic/open population, this tree can instead be analyzed using parallel trials simulation.

As with dynamic cohort models, the dynamic simulation model's preferences are set to allow non-coherent probabilities, as illustrated in the previous chapter.

If this preference is turned on, running a microsimulation will force parallel trials. Additionally, TreeAge Pro will force the “Number of trials” setting to 1 (with additional parallel trial/individuals generated as needed via non-coherent initial and transition probabilities).



Setup parallel dynamic trials

The microsimulation report will show only a *single line of results*. (Use the GlobalIN function to output per-trial reports dynamically.)

Use the additional parallel trials options (*Number of sets*, *Value of each parallel set trial*), shown above, to control whether an average or sum is reported for the group of trials.

The rationale for running multiple sets of parallel trials is that normal, sequential trials have no variance due to interaction. Basically there is no “population” per se, just independent individuals, so we run as many trials as possible to average out their variation. In the case of parallel trials (including dynamic simulations), *trial interactions* are a possible new source of variation – both for individual outcomes and for population outcomes. Each run of a hypothetical population in a disease outbreak simulation, for example, may result in a very different size and pattern of outbreak (for the same model parameters). In an extreme case, for example, the simulation might be highly dependent on what happens to just the first few individuals that encounter an infectious agent.

Additional details on parallel trials are provided below.

- The sum of the *initial probabilities* defines the initial size of the “population.” In the prior example, 10001 trials are created via initial “probabilities”.
- Parallel trials created using initial probabilities are placed in order into the states. In the example, trial #1 starts in the top state, trials #2 to #5001 start in “male” and so on. (As usual, the `_trial` keyword will return this integer index for the currently running trial.)
- Non-coherent transition (or initial) probabilities must sum to an integer. With non-coherent transition probabilities, the current trial follows the first 1.0 of total probability. In the example, at “birth” a “pregnant female” follows the “mother” arm.

- The remainder transition probability – i.e., the sum minus 1.0 – “spawns” new trials at the end of the trials list. In the example, during each cycle, trial #1 spawns 75 new trials (#10002 to #10077 and so on).
- Newly-created trials begin the next cycle at the node after where they were created. Unless `_CLOCK` is being used to synchronize trials (see below), the `_stage` counter increments “normally” – a new trial created during `_stage 0` will start its first cycle in `_stage 1`.
- If the total transition probability = 0, then the trial is terminated, but not deleted, so its final values will count in the expected value output. An alternative might be to “recycle” the individual by resetting trackers, etc.

Parallel trial details

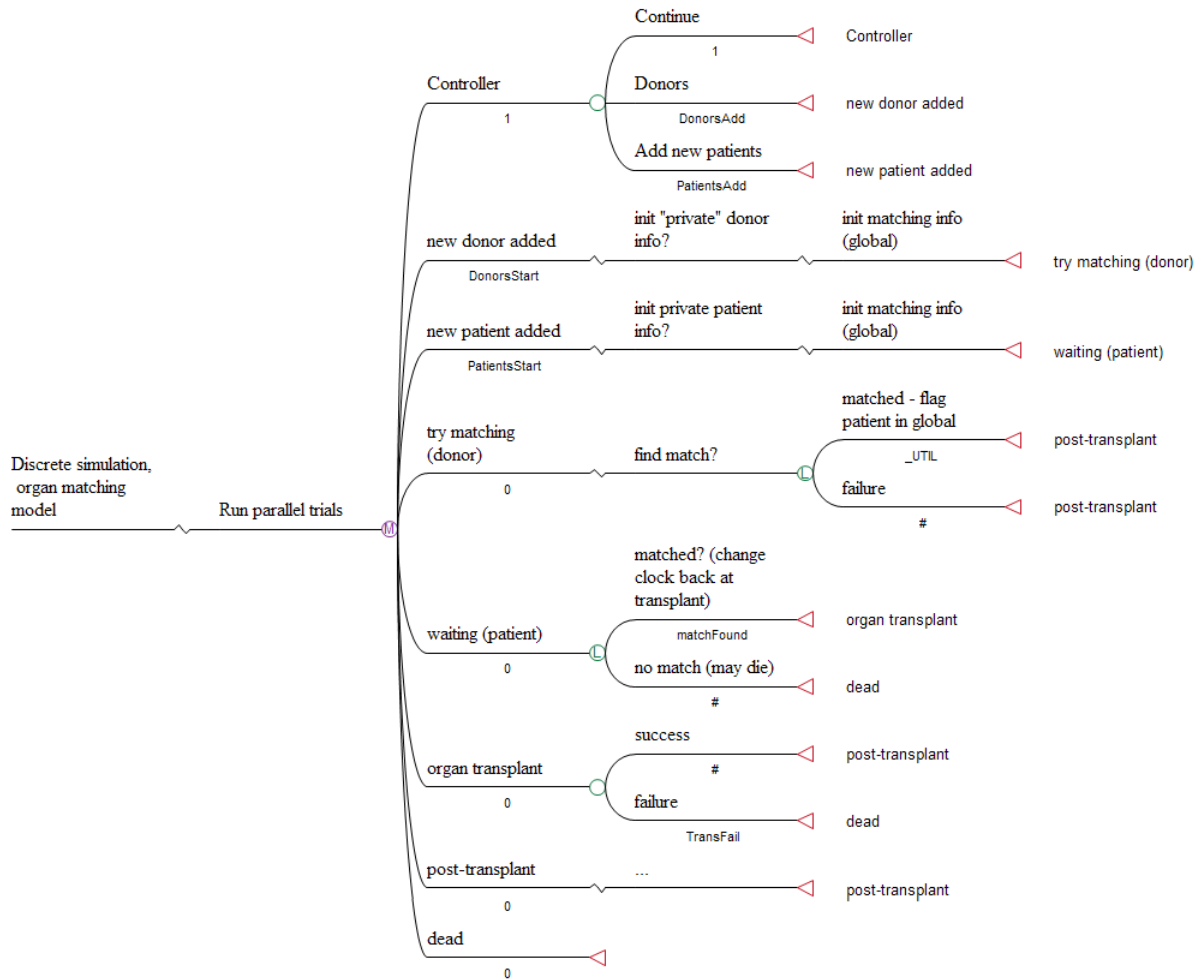
36.8.5 Using `{T} _CLOCK` with dynamic parallel trials

In a dynamic simulation model, the parallel trials are evaluated in an orderly fashion by default, one cycle at a time. The `_stage` counter increments after processing one cycle for all trials that existed at the start of the cycle.

However, if `{T} _CLOCK` exists in the dynamic simulation model, synchronization of parallel trials will occur:

- When non-coherent transition probabilities spawn new trials; the parent trial's `_CLOCK` is copied to the spawned trials. (No other trackers are copied.)
- A newly-created trial stores the integer index of its “parent” trial. The new keyword `_parallel_trial_creator` is used to reference this number.

The Health Care tutorial example tree shown below, “Organ Allocation”, dynamically creates donor and patient trials, along with a `{T} _CLOCK` patient-time tracker. Synchronization of donors and patients is important when dealing with the time dependency of organ transplant supply and demand.



Organ Allocation tree

36.8.6 Global matrices and parallel trials: communication, competition, and reporting

The GlobalIN and Global functions provide a convenient means of handling two complex tasks in microsimulation models, particularly when using dynamic/parallel trials:

- As noted in the introduction to this section, trackers work well for storing private (per-trial) states, but do not handle communication between trials or storage of “public” or system-level states; and
- The level of reporting, available by default, in microsimulation output may be insufficient for some analysis tasks.

Global matrices handle both needs in a very flexible way.

The need for a “public” storage area (e.g., for system-level status, or for communicating values/information between trials) is a requirement of some discrete event simulation models. For example, competition for limited resources may need to be modeled.

The “Organ Allocation” example model in the previous section makes extensive use of the `GlobalN()` function to store and retrieve system-level information regarding the availability (and matching characteristics) of organ donors. This global information is not only available during the simulation, but is exported (in this case, to Excel worksheets) after the simulation completes.

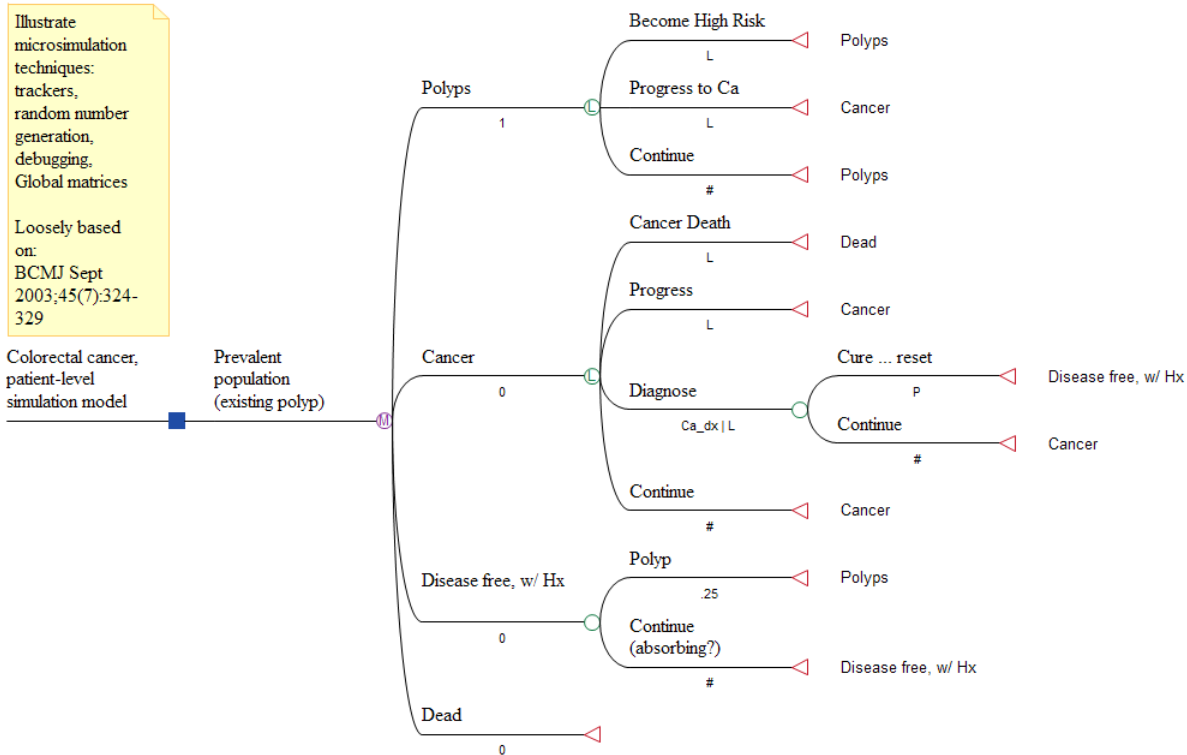
The example also makes use of a Python user-defined function to search for matches between the patient and donor Global matrices.

The Global matrices have a wide variety of uses in reporting and debugging. For example, in a simple microsimulation (no PSA outer loop), tracker variables report only their final value for each trial. Reports do not provide intermediate values of trackers, although that may be of interest. Also, simulation does not report changing state probabilities over time (as in the cohort analysis State Probabilities graph).

In the context of a dynamic parallel trials simulation, a further reporting issue is the fact that only a single line of output is provided in the simulation output (providing either the average or sum of the list of trials). A similar issue exists in the output of 2-dimensional probabilistic sensitivity analysis utilizing microsimulation in the inner loop.

Whereas tracker variables are not designed to handle such complex or fine-grain reporting tasks, the Global matrix-related functions are intended for just such purposes. Global matrices can be used to record changing trial state, tracker values, or other information per-stage into separate rows, columns, or N matrices for any/all sample iterations). Records can even be kept for multiple samples in a PSA loop.

The Health Care tutorial example tree shown below, “Cancer Simulation”, utilizes `GlobalN` and related functions to store a high level of detail about each trial.



Cancer Simulation tree

Although not a dynamic/parallel trials model, the same GlobalN techniques used in the example can be applied to parallel trials models.

36.8.7 Additional parallel trials features

New built-in keywords supporting parallel trials include:

`_parallel_trials_sets_size` (# of sets in loop)

`_parallel_trials_set` (# of current set)

Python user-defined functions (see Chapter 21) can create a parallel trials object which allows one trial to interact with another without using a Global matrix:

`para = treeage.getParallelTrials()` (returns an object with access to parallel trials)

`para.getTrialN_Clock(int trial)` (returns the `_CLOCK` value for the specified trial number)

`para.setTrialN_Clock(int trial, double v)` (sets the `_CLOCK` value for the specified trial number)

`para.getTrialNTracker(int trial, string var)` (returns a tracker value for the specified trial number)

`para.setTrialNTracker(int trial, string var)` (sets a tracker value for the specified trial number)

36.9 2-dimensional probabilistic sensitivity analysis using microsimulation

Some Markov models require microsimulation, either for tracker variables or per-trial sampling distributions. Performing sensitivity analysis on these microsimulation models will generally require probabilistic sensitivity analysis using Monte Carlo simulation (rather than n-way, expected value analyses).

The probabilistic sensitivity analysis method samples sets of parameter values and, for each set, recalculates the expected values for the model. The Monte Carlo Simulation, Distributions and Probabilistic Sensitivity Analysis Chapter covers the general usage of Monte Carlo simulation to perform probabilistic sensitivity analysis.

The theoretical background for this kind of probabilistic sensitivity analysis is discussed in detail elsewhere, for example, in papers published in the Journal of the Society for Medical Decision Making including:

“Monte Carlo Probabilistic Sensitivity Analysis for Patient Level Simulation Models,” Anthony O'Hagan, et al, *Health Economics*, 16:1009–1023 (2007).

“Representing First- and Second-order Uncertainties by Monte Carlo Simulation for Groups of Patients,” Elkan Halpern, Milton Weinstein, Maria Hunink, G Scott Gazelle, *Med Decis Making* 20:314-322 (2000).

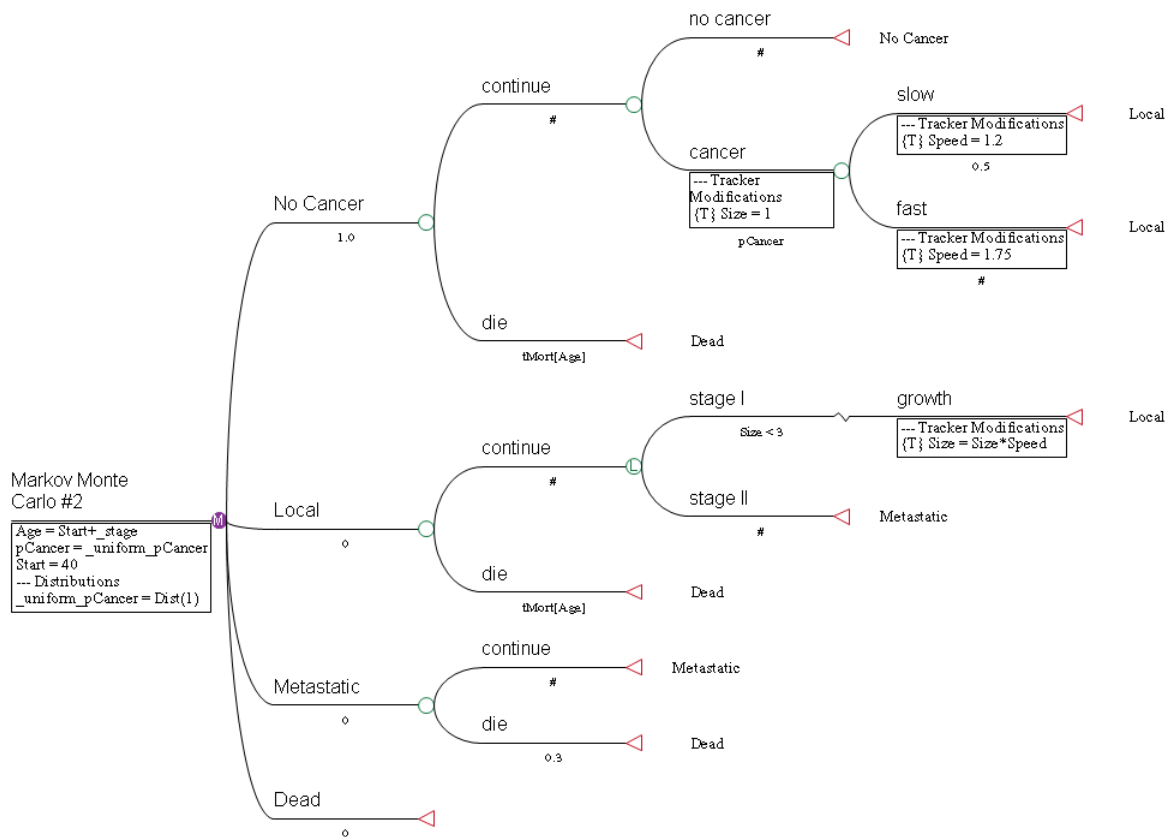
You are urged to explore these and other publications on this topic.



The tracker values reported for 2-dimensional simulations, which sample distributions and then run microsimulation trials for each set of sample values, are averages for each group of trials run for a particular set of parameter samples.

36.9.1 Performing sensitivity analysis with microsimulation

The Health Care tutorial example model shown below – “Markov Monte Carlo #2” – uses tracker variables in Markov calculations, specifically the logic node branch expression at Local > continue > stage I. This means that simulation trials must be used to correctly analyze the model.



Markov Monte Carlo #2 tree

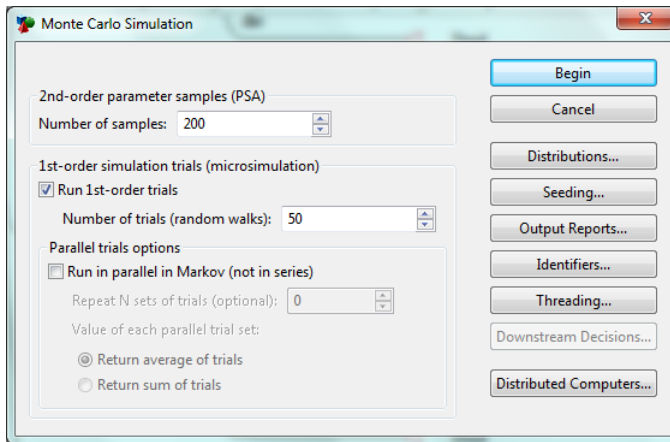


This model requires a table, tMort[], created in the previous chapter. Create the table now. If you have already done so, you can export/import the table from the model created in the previous chapter via a global tables file.

Since expected value calculations are not valid in the model, probabilistic sensitivity analysis must be used to perform uncertainty analysis (let's say on pCancer, in this case).

To perform a probabilistic sensitivity analysis using groups of individual trials:

- In the Markov Monte Carlo #2 tree, select the root node.
- Choose Analysis > Monte Carlo Simulation > Sampling + Trials.
- In the Monte Carlo Simulation dialog, specify 200 distribution samples and 50 trials per sample, and then press enter or click Begin.



Two-dimensional simulation setup

36.9.2 Interpreting a probabilistic sensitivity analysis using microsimulation

The final output of the simulation will report the statistical summary for the analysis. The statistics are calculated as follows:

1. Each group of trials is averaged to produce a mean value which represents that set of distribution samples. This averaging is applied to all output quantities – not just the tree's cost and/or effectiveness attributes, but tracker values and reported distribution sample values, as well.
2. The resulting sets of average values are then statistically analyzed for probabilistic sensitivity analysis.

If you click on the "Values, Dists, Trackers" link to the right of the simulation statistics, TreeAge Pro will display a list of averages, each simulating an expected value calculation by averaging the results of multiple trials. For each iteration (i.e., sample), average values are also reported for tracker values and distribution samples, again based on that iteration's group of trials.

In essence, a mean value from a group of trials is estimating expected value (EV) for the model. Within the context of probabilistic sensitivity analysis (PSA), this EV estimate is used the same way as the EV calculations from PSA described in the Monte Carlo Simulation, Distributions and Probabilistic Sensitivity Analysis Chapter. All outputs are the same.

36.9.3 How many samples and trials?

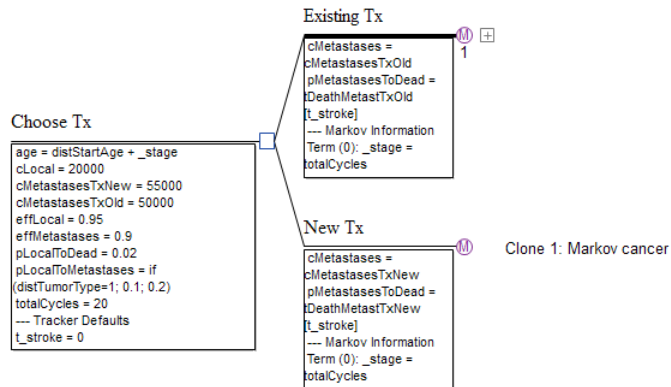
The number of samples and trials required for stable, accurate results depends greatly on the complexity of the model. For a helpful discussion of determining the number of samples (outer loop= N) and trials (inner, microsimulation loop= n) required to support your analysis objectives, refer to:

- "Monte Carlo Probabilistic Sensitivity Analysis for Patient Level Simulation Models," Anthony O'Hagan, et al, *Health Economics*, 16:1009–1023 (2007).

36.10 Sensitivity Analysis and Microsimulation

In prior versions of TreeAge Pro, it was not possible to run a one-way sensitivity analysis and Microsimulation at the same time. However, TreeAge Pro 2012 introduced this analysis option.

Open the Health Care tutorial example model "Microsimulation Markov Clone."

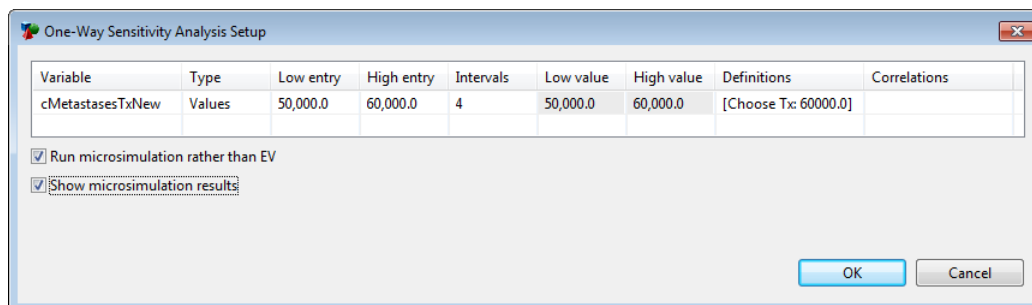


Microsimulation Markov Clone model

This model requires Microsimulation because it uses individual trial-level distributions and trackers. However, we also want to run one-way sensitivity analysis on the variable *cMetastasesTxNew*.

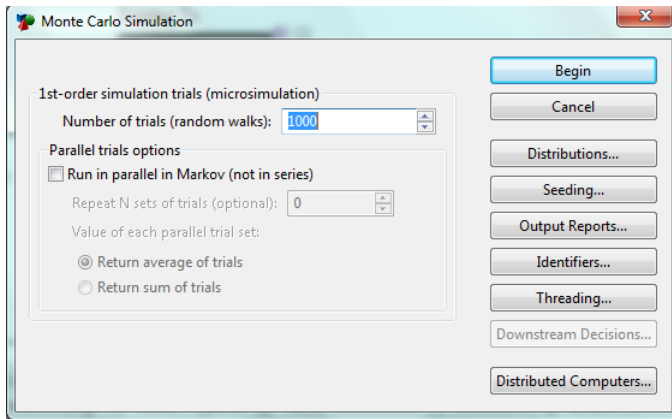
To run one-way sensitivity analysis:

- Select the root node.
- Choose Analysis > Sensitivity Analysis > 1 Way.
- Choose the variable *cMetastasesTxNew* and enter the low, high and interval values as seen below.
- Select the option "Run microsimulation rather than EV".
- Select the option "Show microsimulation results".
- Click OK.



One-Way Sensitivity Analysis Setup Dialog

The two options we selected are related to Microsimulation. The first initiates the Microsimulation, while the second displays the individual Microsimulation results as well as the sensitivity analysis results. Since we chose to run Microsimulation, the appropriate simulation dialog is then presented.



Monte Carlo Simulation Dialog for Microsimulation

All the options available in this dialog are described in a previous section of this chapter. Click Begin.

Based on the sensitivity analysis options, the analysis needs to consider values of \$50K, \$52.5K, \$55K, \$57.5K and \$60K for variable *cMetastasesTxNew*. In a regular sensitivity analysis, expected values would be generated for each variable value using Markov Cohort Analysis. However, since we submitted Microsimulation within the sensitivity analysis, five separate sets of trials are executed, one for each variable value. Since we asked to show the Microsimulation results, we will see those results as well as the overall sensitivity analysis results.

Monte Carlo C-E Statistics							
Attribute	Statistic	Existing Tx	New Tx				
Cost	Mean	316,785.00	363,985.00				
	Std Deviation	18,820.07	209,130.63				
	Minimum	10,000.00	10,000.00				
	2.5%	30,000.00	30,000.00				
	10%	80,000.00	90,000.00				
	Median	310,000.00	370,000.00				
	90%	580,000.00	650,000.00				
	97.5%	745,000.00	805,000.00				
	Maximum	930,000.00	985,000.00				
	Sum (n*Mean)	316,785,000.00	363,985,000.00				
	Size (n)	1,000.00	1,000.00				
	Variance	34,901,738,775.00	44,028,894,775.00				
	Variance/Size	34,901,738.77	44,028,894.77				
	SQRT(Variance/Size)	5,907.77	6,635.43				
Eff	Mean	9.39	10.24				
	Std Deviation	5.54	5.61				

Sensitivity Cost Effectiveness Analysis							
cMetastasesTxNew	Strategy	Cost	Incr cost	Eff	Incr Eff	C/E	Incr C/E (ICER)
50000.0	Existing Tx	316785.0	0.0	9.3885	0.0	33741.99149	0.0
	New Tx	363985.0	47200.0	10.23805	0.8496	35552.18035	55555.55556
52500.0	Existing Tx	315180.0	0.0	9.3926	0.0	33556.20382	0.0
	New Tx	375323.75	60143.75	10.26155	0.86895	36575.73661	69214.2816
55000.0	Existing Tx	313180.0	0.0	9.3094	0.0	33641.26582	0.0
	New Tx	374207.5	61027.5	9.99925	0.68985	37423.55677	88464.88367
57500.0	Existing Tx	312835.0	0.0	9.33357	0.0	33517.16786	0.0
	New Tx	400992.5	88157.5	10.27362	0.94005	39031.25722	93779.58619
60000.0	Existing Tx	313940.0	0.0	9.14195	0.0	34340.59473	0.0
	New Tx	404870.0	90930.0	9.9191	0.77715	40817.21124	117004.4393

Sensitivity Analysis and Microsimulation Results

The mean values from the first set of Microsimulation results are used for the first set of sensitivity analysis EV results as shown above. The other four sets of Microsimulation mean values are used for the remaining sensitivity analysis EVs. This leaves you with a standard sensitivity analysis output as described in earlier chapters (simple, cost-effectiveness).



As the sequence runs individual Microsimulations, the variable definition in the model is changed. You will need to change it back to the original value after the last simulation.

36.11 Sequences and linear sensitivity analysis

In TreeAge Pro, it is possible to create sequences of repeated analyses, including Monte Carlo simulations, as described in the Stored Analysis Abstracts and Sequences Chapter. One potential application of the analysis sequencing feature is in creating a linear series of Monte Carlo simulations.

The sequenced simulations approach allows point values to be specified for a parameter and then a microsimulation to be performed in an automated fashion. This results in something similar to TreeAge Pro's standard n-way sensitivity analysis.

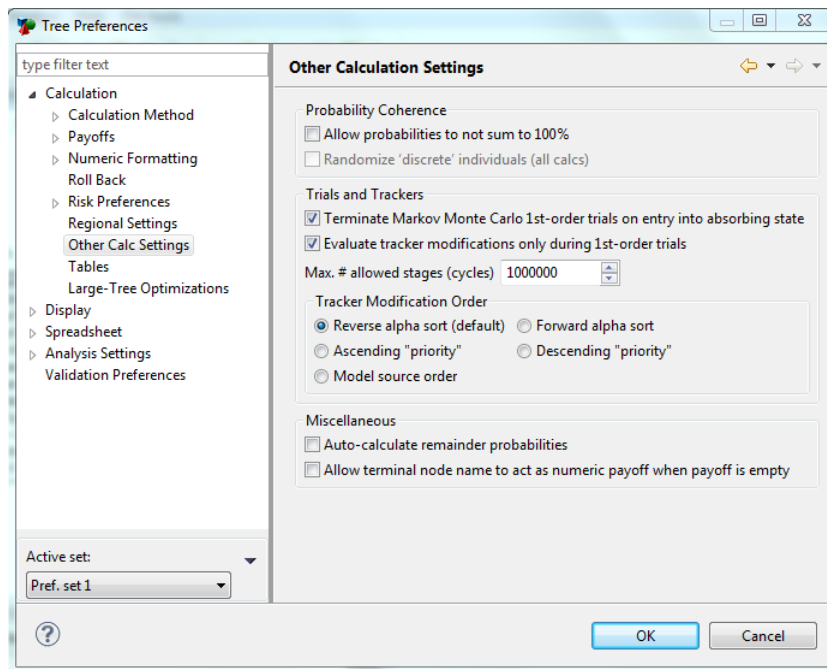
The analysis sequencing feature first requires that a “template” simulation be created as a stored analysis, to serve as the instructions to TreeAge Pro for each simulation in the sensitivity analysis sequence. The final output of the sequence will not be a sensitivity analysis line graph, but a series of simulation output windows. In order to be able to differentiate between the simulation output windows for the various values of the variable (e.g., pCancer), you will use identifying values in the initial simulation. This way, the value of pCancer will be reported in each simulation output window and its text report. You could use the output from the simulations to generate a sensitivity analysis graph in external software.

36.12 Other aspects of microsimulation

Markov microsimulation is a very broad topic and TreeAge Pro includes numerous microsimulation options and features which may be of interest in select instances.

36.12.1 Markov termination during simulation trials

By default, Markov process calculations terminate during microsimulation trials when *either* the termination condition is true *or* a trial enters an absorbing state. It is possible to turn off the latter behavior so that trials will continue processing even after entry into an absorbing state. Changing this default setting is likely to be appropriate in only a few Markov models (e.g., where absorbing states do not correspond to dead states or other endings of the Markov process). This setting is found in the Other Calculation Settings category of the Preferences dialog. The default for this option is checked.



Tree Preferences - Terminate trials

The Terminate upon entry into absorbing state setting generally substitutes for the threshold portion of the default Markov termination condition (i.e., “_stage_eff < 0.001”). The threshold part of the condition should generally be removed for microsimulation models.

36.12.2 Using logic nodes in a Markov microsimulation

A *logic node* acts like a decision node, in that it selects one path from its branches; rather than looking at expected values, however, it chooses a path by evaluating logical expressions. Starting at the top branch, the first node with an expression that evaluates to *true* is selected. A simple logic node might have two branches, X and Y, with the expression `_stage > 4` below branch X and `_stage <=4` (or `#`) below branch Y. When the logic node is encountered, either branch X or branch Y is followed based on the current value of `_stage`.

Most logical expressions in Markov processes will reference the values of tracker variables. For example, in Markov Monte Carlo #2 and #3, a tracker variable serves to remember the current size of a tumor during a simulation trial, while a logic node determines whether to transition to a Metastatic state based on the value of this tracker.

37. Markov Technical Details

This chapter has not yet been written.

38. Time-to-Event Simulation

The Individual-Level Simulation and Markov Models Chapter described how to run individuals through a Markov model. This chapter focuses on time-to-event models, which extend concepts further by allowing you to structure models around events and continuous time rather than health states and time cycles of fixed length.

38.1 Time-to-event models

Time-to-event models differ significantly from Markov models in that they focus on events rather than health states. TreeAge Pro uses a similar decision tree model structure to Markov models, but uses new node types and keywords. It is important to consider that flow through the model is based on events and the progression of continuous time and not discrete time intervals.

Time-to-event models contain most of the elements found in Discrete Event Simulation (DES) models. However, TreeAge Pro time-to-event models do not directly support resource constraints, queues and other elements less frequently used for healthcare models. Within this document, we will use DES and time-to-event somewhat interchangeably.

Similar to Markov models, time-to-event models can be incorporated into a large decision tree to facilitate cost-effectiveness analysis and other mechanisms to compare treatment strategies. Unlike Markov models, time-to-event models can only be executed via Microsimulation, which sends individual patients through the model and generates expected values by aggregating the individual patient histories into a mean.

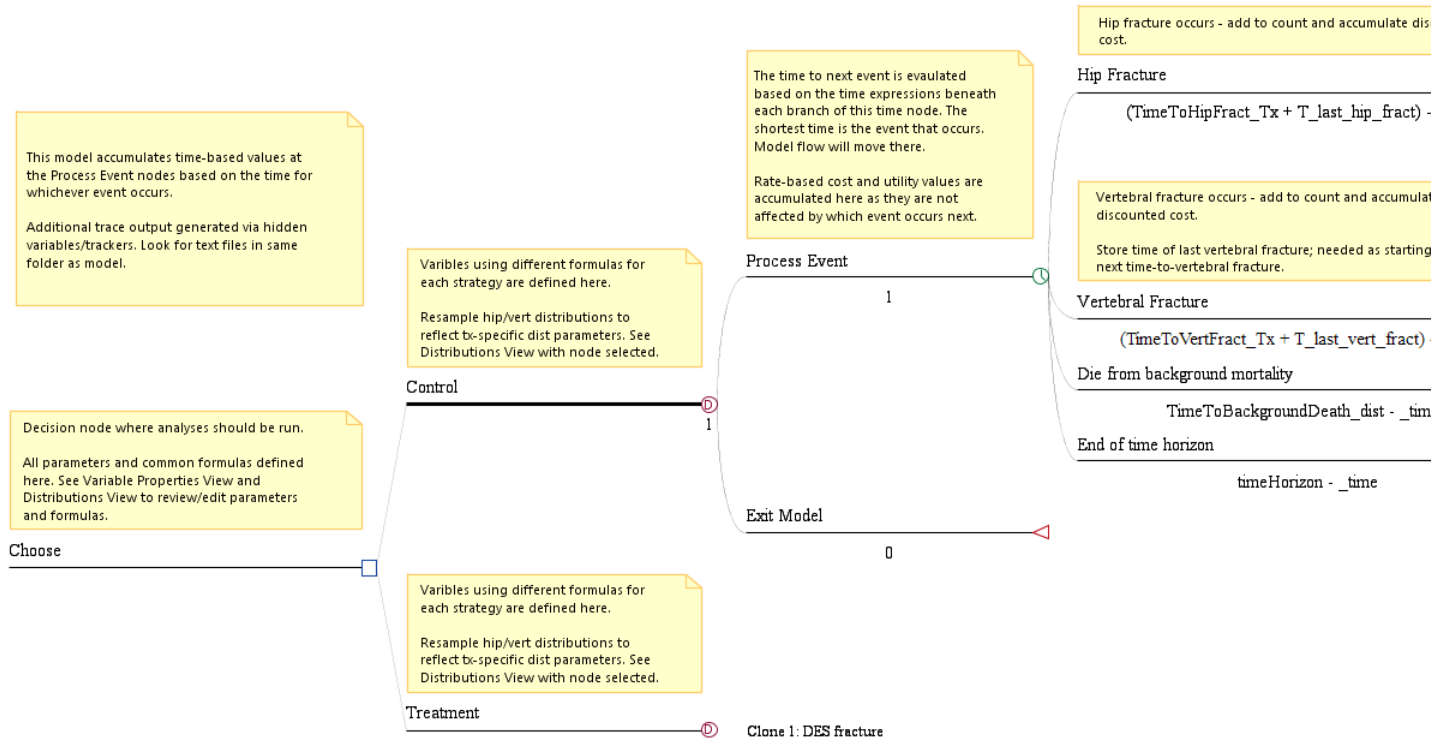


Time-to-event models must be analyzed via Microsimulation to send individual patients through the model. Cohort-level analyses are not supported.

This chapter will use the DES tutorial example model, Osteo DES Model. This model follows three events through a time horizon - hip fractures, vertebral fractures and death.

38.2 Time-to-event model structure

The DES tutorial example tree, Osteo DES Model, is a typical time-to-event model. Note that variables are hidden in the figure below.



Time-to-event model structure

The model has two strategies, each of which starts with a *DES node*. The DES node then has two branches - a *time node* for the process event loop and a terminal node to exit the model. To the right of the Process Event node, there are four possible events, two which return to the process event loop and two which exit the model.

This type of process event loop is a typical way to structure a time-to event model. The remainder of this chapter will highlight the critical elements within this model.

38.3 DES Keywords

Time is a critical element in time-to-event models. TreeAge Pro provides keywords to help track time within the model. Additionally, new keywords were added to access rewards (payoffs) accumulated by a patient during simulation (total and by "cycle").

Keyword	Description
<code>_time</code>	Current time for the patient.
<code>_event_time</code>	Time of the last event.
<code>_prior_time</code>	Time before the last event.
<code>_total_rwdx[index]</code>	Total value accumulated by patient in reward number <i>index</i> from beginning of time to current event.

Keyword	Description
<code>_stage_rwdx[index]</code>	Value accumulated by patient in reward number <i>index</i> from beginning of cycle (path from DES node to current event).

DES keywords

The `_time` keyword is the most commonly used keyword within time-to-event models. It is often used in the following ways.

- Discounting rewards.
- Calculating a time-to-event from current time forward.
- Storing the time of an event.
- Tracing patient history.

The `_event_time` keyword is used less frequently, but it can be useful for tracing patient history.

The `_prior_time` keyword is used for discounting time-based rewards from the prior event to the current event. This is described in detail later in this chapter.

38.4 DES Node Type

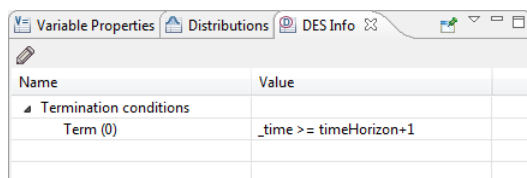
Time-to-event models include two new node types - DES and Time.



Time-to-event node types - DES and Time

The DES node type indicates the beginning of the time-to-event model, similar to a Markov node beginning the Markov model. The only time-to-event element provided at this node is the termination condition, which indicates when the analysis should stop. For DES models, this would typically be a time horizon.

All DES-related information is entered via the DES Info View. Similar to the Markov Info View, the contents of this view change based on the context within the DES model. At a DES node, the termination condition is entered.



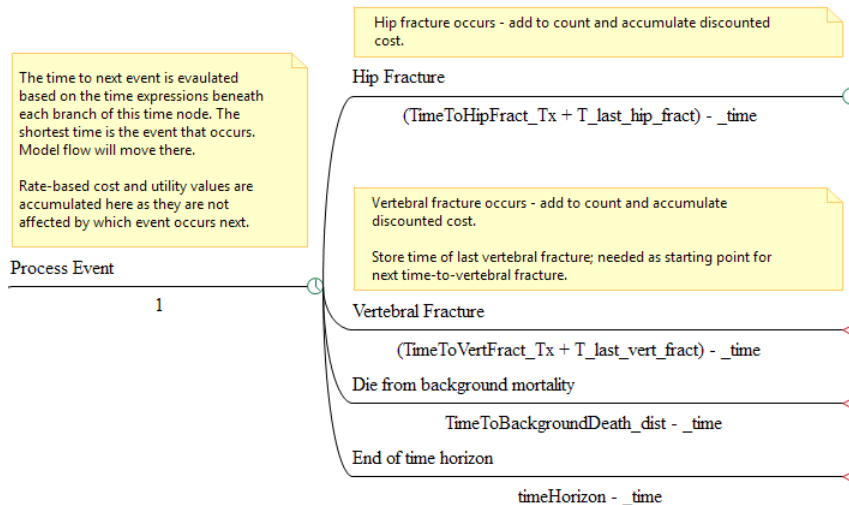
DES Info View with DES node selected

In the Osteo DES Model, the termination condition is true when the patient `_time` exceeds the `timeHorizon` variable (25 years) + 1, processing of the patient will stop, even if the patient has not reached the node Exit Model. However, all patients should reach the Exit Model node.

38.5 Time Node Type

Time nodes are the key to time-to-event models. Time nodes look similar to chance nodes in that there are expressions under each branch node. However, each expression represents the time-to-event for that event. The patient will move from the time node to the event with the shortest time-to-event.

Consider the time node in the Osteo DES Model as shown below.



Time node and events

Note that each event (the two fracture types, background mortality and end of time horizon) has a time expression beneath it.

The *Hip Fracture* time-to-event includes three elements.

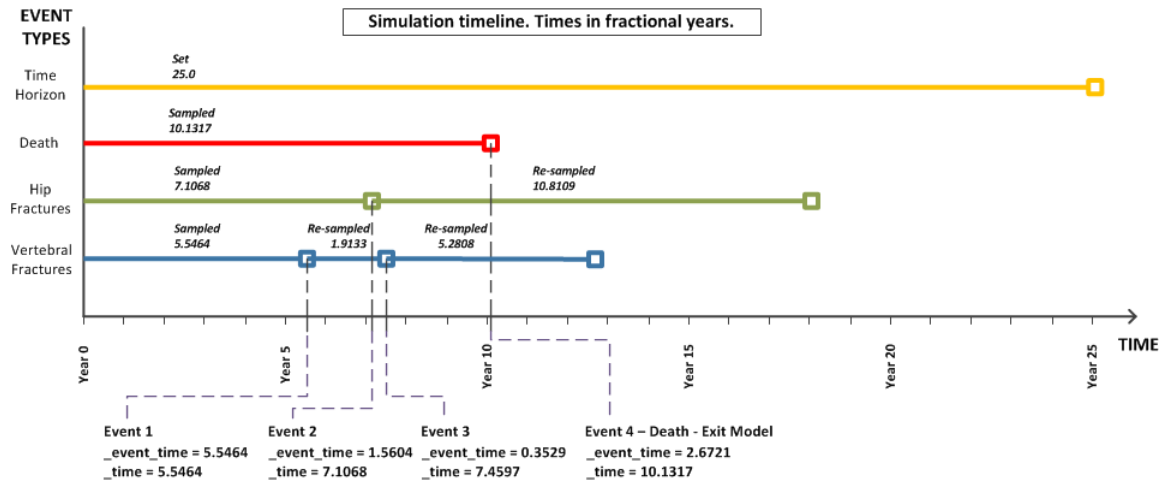
- *TimeToHipFract_Tx* references the appropriate Weibull distribution for the *Control* or *Intervention* strategy.
- *T_last_hip_fracture* is the time of the last hip fracture. The new hip fracture time starts after the last hip fracture, so this must be included in the time-to-event.
- *_time* is subtracted from the time-to-event to reflect that some time may already have passed since time 0.

As a whole the *Hip Fracture* time-to-event is the time to the next fracture, starting with the time of the last fracture, less the time that has already passed. The *Vertebral Fracture* time-to-event functions in a similar way, but with different distributions and a different tracker.

The *Die from background mortality* time-to-event references the time-to-death distribution directly since there is no difference between the strategies. Again, the current time is subtracted via the *_time* keyword.

The *End of time horizon* time-to-event references the total *timeHorizon* variable (25 years) less the current time.

The following timeline examines the flow of a specific patient as he/she progresses through the model in the *Control* strategy.



Patient event timeline

Here is the basic patient progression:

- As the patient enters the model, all distributions are sampled for the next time-to-event. The samples are hip fracture at time 7.1068, vertebral fracture at time 5.5464 and death at time 10.1317.
- The shortest time is for vertebral fracture, so that event occurs at time 5.5464.
- The vertebral fracture distribution is resampled, drawing a value of 1.9133. Added to the prior vertebral fracture time, the next vertebral fracture may occur at time 7.4597.
- The next event is a hip fracture, which occurs at time 7.1068.
- The hip fracture distribution is resampled, drawing a value of 10.8109. Added to the prior hip fracture time, the next hip fracture may occur at time 17.9177.
- The next event is the second vertebral fracture at time 7.4597.
- The vertebral fracture distribution is resampled, drawing a value of 5.2808. Added to the prior vertebral fracture time, the next vertebral fracture may occur at time 12.7405.
- The next event is death at time 10.1317, and the patient leaves the model.

Patient progression through model

38.6 Accumulating value - rate and fixed rewards

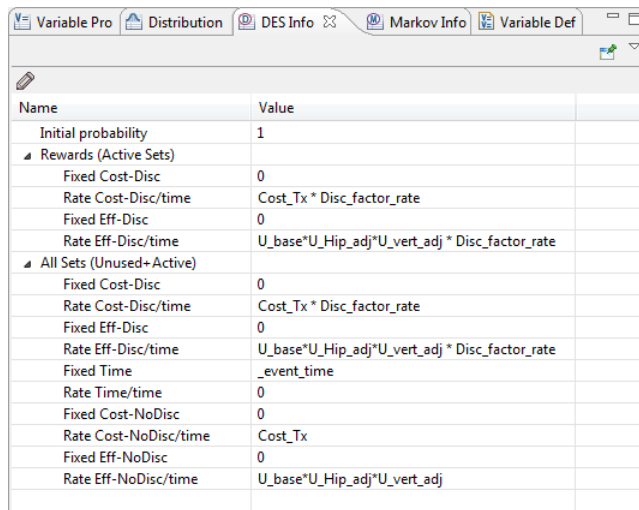
In Markov models, value is accumulated via state rewards (cycle in health state) and transition rewards (events). In DES nodes, the equivalents are rate rewards and fixed rewards.

38.6.1 DES rate rewards

Markov state rewards are accumulated based on the patient being in a health state for a fixed period of time. In DES nodes, there are no fixed cycles. Therefore, the amount of value that is accumulated before an event needs to be calculated based on a rate reward and the time-to-event.

Each time an event occurs, the *_event_time* is automatically calculated. Rate rewards are multiplied by the *_event_time* automatically to calculate the value accumulated at a specific rate for the time to the event. Let's examine this within the context of the Osteo DES Model.

The figure below contains the DES Info View with the *Process Event* node selected.



Name	Value
Initial probability	1
▲ Rewards (Active Sets)	
Fixed Cost-Disc	0
Rate Cost-Disc/time	Cost_Tx * Disc_factor_rate
Fixed Eff-Disc	0
Rate Eff-Disc/time	U_base*U_Hip_adj*U_vert_adj * Disc_factor_rate
▲ All Sets (Unused+Active)	
Fixed Cost-Disc	0
Rate Cost-Disc/time	Cost_Tx * Disc_factor_rate
Fixed Eff-Disc	0
Rate Eff-Disc/time	U_base*U_Hip_adj*U_vert_adj * Disc_factor_rate
Fixed Time	_event_time
Rate Time/time	0
Fixed Cost-NoDisc	0
Rate Cost-NoDisc/time	Cost_Tx
Fixed Eff-NoDisc	0
Rate Eff-NoDisc/time	U_base*U_Hip_adj*U_vert_adj

DES Info View with Time node selected

Note that the active payoffs (1 & 2) are discounted cost and effectiveness, respectively. Additional payoffs are enabled. Let's examine the Cost-NoDisc and Eff-NoDisc (both not discounted) payoffs for now. Discounting will be described later in this chapter.

Note the value *Cost_Tx* as the value for undiscounted cost and a utility formula as the value for undiscounted effectiveness. When the model runs, the time node will select the next time-to-event based on the shortest time-to-event, resulting in the *_event_time*.

Let's assume that the following time-to-event values are calculated from formulas/distributions...

- (TimeToHipFract_Tx + T_last_hip_fract) - _time = 9.388511241
- (TimeToVertFract_Tx + T_last_vert_fract) - _time = 3.744567566
- TimeToBackgroundDeath_dist - _time = 7.072967153
- timeHorizon - _time = 25.0

Time-to-event values for time node

The shortest time is the time-to-vertebral fracture, so that event occurs. At the *Process Event* node, it does not matter which event occurs for value accumulation, just the time-to-event.

The non-discounted effectiveness is calculated by evaluating the expression entered and multiplying it by the time-to-event.

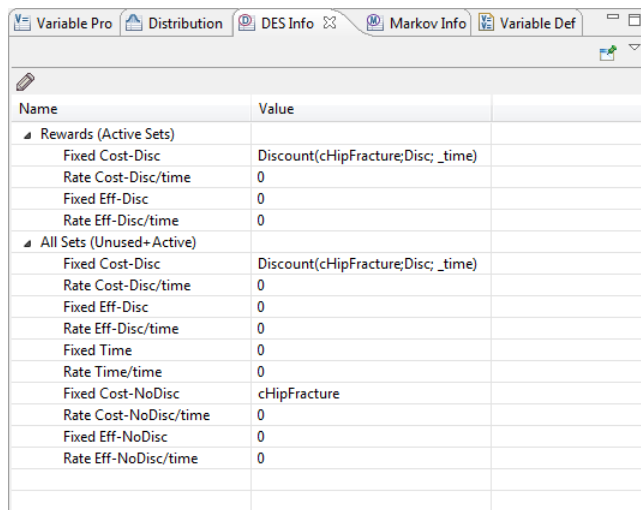
$$(U_base * U_hip_adj * U_vert_adj) * time_to_event = (0.7) * (3.744567566) = 2.62119729628424$$

Therefore, 2.62119729628424 QAL Ys were accumulated by the time the vertebral fracture occurred.

38.6.2 DES fixed rewards

Generally, event-driven rewards are fixed rather than time-based. Therefore, event rewards are placed in the fixed reward entries via the DES Info View.

The figure below contains the DES Info View with the *Hip Fracture* node selected.



Name	Value
▲ Rewards (Active Sets)	
Fixed Cost-Disc	Discount(cHipFracture;Disc;_time)
Rate Cost-Disc/time	0
Fixed Eff-Disc	0
Rate Eff-Disc/time	0
▲ All Sets (Unused+Active)	
Fixed Cost-Disc	Discount(cHipFracture;Disc;_time)
Rate Cost-Disc/time	0
Fixed Eff-Disc	0
Rate Eff-Disc/time	0
Fixed Time	0
Rate Time/time	0
Fixed Cost-NoDisc	cHipFracture
Rate Cost-NoDisc/time	0
Fixed Eff-NoDisc	0
Rate Eff-NoDisc/time	0

DES Info View with event node selected

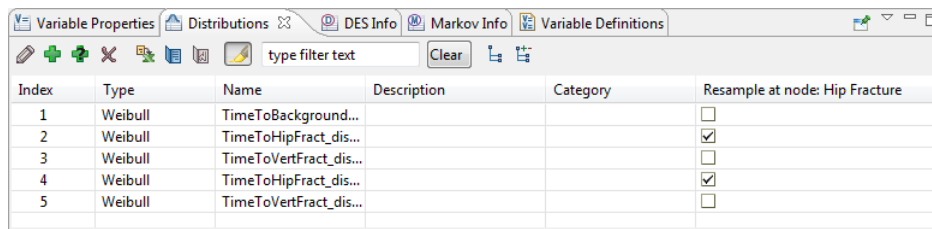
Focusing on the undiscounted cost, the cost of the hip fracture (parameter variable *cHipFracture*) is placed in the fixed reward rather than the rate reward.

38.7 Distribution Sampling

In most ways, distributions work the same as in other types of decision trees as described in the Monte Carlo Simulation, Distributions and Probabilistic Sensitivity Analysis Chapter (for PSA/parameter sampling) and the Individual-Level Simulation and Markov Models Chapter (for Microsimulation/individual characteristics). However, time-to-event simulations require additional control over distribution sampling beyond the standard sampling rates.

Consider the time-to-event distribution samples in the Osteo DES Model for hip fracture. When a patient starts its path through the model, the time-to-hip fracture distribution is sampled based on the sampling rate of "Once per trial". However, when a hip fracture occurs, a new sample is needed for hip fracture, so that the identical time is not used.

TreeAge Pro allows for distribution sampling at any node via the Distributions View.



Index	Type	Name	Description	Category	Resample at node: Hip Fracture
1	Weibull	TimeToBackground...			<input type="checkbox"/>
2	Weibull	TimeToHipFract_dis...			<input checked="" type="checkbox"/>
3	Weibull	TimeToVertFract_dis...			<input type="checkbox"/>
4	Weibull	TimeToHipFract_dis...			<input checked="" type="checkbox"/>
5	Weibull	TimeToVertFract_dis...			<input type="checkbox"/>

Distributions View - resampling

When a patient reaches the Hip Fracture node, the hip fracture event occurs, and a new time-to-hip fracture sample must be drawn. Note that there are two hip fracture distributions, one for each strategy. Because the primary event loop is cloned, either distribution could be used, so both are resampled.



Notes on distribution resampling:

Unlike variables, distributions are designed with global scope. Therefore, distribution parameters can only be defined within the distribution and/or by referencing the root node definitions for variables. If you redefine a variable to the right of the root node, that definition will not be used when the distribution is resampled.

However, expressions used in distribution parameters (including root node variable definitions) are calculated dynamically, so the parameter values can change as a patient progresses through the model.

For example, if a distribution parameter references a tracker or keyword (i.e., `_time`), the current values will be used when the distribution is resampled.

38.8 Tracking Events

In time-to-event models, events can be tracked via trackers or in reward sets. While these two storage mechanisms are quite different in a Markov context, they function quite similarly in time-to-event models. Full description of trackers and tracker views is available in the Individual-Level Simulation and Markov Models Chapter.

In the Osteo DES Model, trackers are used to count fractures as well as record the time of the last fracture. This use does not differ from the use in any Microsimulation model. Trackers are often used in this way because it is easy to reference the current tracker values as needed in the model. For example, the time of last fracture is needed in calculating the time to the next fracture.

Rewards in the DES context can also be stored and retrieved at any time. Rewards are accumulated as described in the "Accumulating value" section of this chapter. The current value of that reward is available via the `_total_rwdx[index]` keyword described earlier, where the index refers to the number of the reward set.

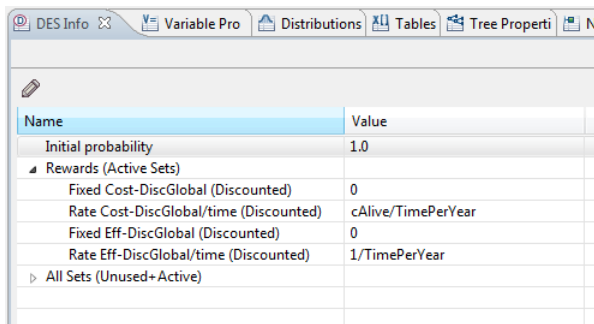
38.9 Discounting in DES Models

In the context of Markov modeling, discounting is far simpler. State rewards are accumulated at specific times in the overall sequence of events. The Discount function calculates the discounted value based on an interest rate and the fixed time. The Discounting rewards section provides more information about how to do discounting with Markov Models. The Discount function and its underlying formula is described in the Building Formulas Using Variables and Functions Chapter.

Discounting nearly always needs to be applied to an entire fixed reward set. TreeAge Pro has two options for discounting.

1. *Apply a single discount rate to an entire reward set:* Use the Tree Preferences to set a discount rate to be applied to any given reward set. We call this Global Discounting.
2. *Adjust each reward manually:* Use the Discount() function as described further below to change the value of each reward.

To *apply a single discount rate to the entire reward set*, we enter the details in the Tree Preferences as described in the Payoff Preferences section. When Global Discounting is applied to a reward set the word "Discounted" appears after the reward name in the DES Info View, as in the figure below.



Name	Value
Initial probability	1.0
▲ Rewards (Active Sets)	
Fixed Cost-DiscGlobal (Discounted)	0
Rate Cost-DiscGlobal/time (Discounted)	cAlive/TimePerYear
Fixed Eff-DiscGlobal (Discounted)	0
Rate Eff-DiscGlobal/time (Discounted)	1/TimePerYear
► All Sets (Unused+Active)	

Global discounting applied to both Fixed and Rate reward sets, indicated by the "Discounting" label after the rewards name in the DES Info View.

The DES tutorial example model DES.Discounting.trex demonstrates global discounting in a DES model along with use of the DiscountContDES and Discount functions for time-based rewards and fixed rewards respectively. Note that analysis of the model using each discounting option yields the same results.

If you prefer to *adjust each reward manually* within a time-to-event context, discount functions handle fixed event costs using the `_time` keyword for the time of the event.

However, within the time-to-event framework, the values accrued since the last event are calculated when the next event occurs. If these values have been accrued at a constant rate between times *a* and *b*, then the discounted value can be calculated as follows;

$$\text{Discounted Value} = V_{ab} \frac{(e^{[b(-DR_i)]} - e^{[a(-DR_i)]})}{(-DR_i)}$$

a is the start time for the value accrual

b is the end time for the value accrual

V_{ab} is the value accrued at a constant rate from time a to b

DR_i is the instantaneous equivalent of the annual discount rate (DR) given by the following formula;

$$DR_i = \ln(1 + DR).$$

Discounting formula for time-based DES rewards

Note that the discounted value formula above includes the accumulation of time rather than just adjusting for discounting.

TreeAge Pro includes functions to handle continuous-time discounting.

DiscountCont(value; annual_disc_rate; time_start; time_end)

DiscountContDES(value; annual_disc_rate)

Using the formulas above, the first function will discount the *value* parameter based on the *annual_disc_rate* parameter from a start time *time_start* to an end time *time_end*. The second function populates the start and end time parameters with the appropriate values for time-based rewards within a DES model - *time_start* is set to the keyword *_prior_time*, and *time_end* is set to the keyword *_time*. We recommend using the second function within your DES models.



Do not use the DiscountContDES function unless your time is measured in years. If not, you will need to use the DiscountCont function and set the *time_start* parameter to *_prior_time* adjusted for the time measurement (divide by 12 for monthly) and set the *time_end* parameter to *_time* with the same adjustment.

In the Osteo DES Model, there are also a few variables that handle discounting of time-based events using the formulas above in a more manual manner.

$Disc = 0.035$

$Disc_i = \ln(1+Disc)$

$Disc_factor_time = (exp_time*(-Disc_i)) - exp_prior_time*(-Disc_i))/(-Disc_i)$

$Disc_factor_rate = Disc_factor_time / _event_time$

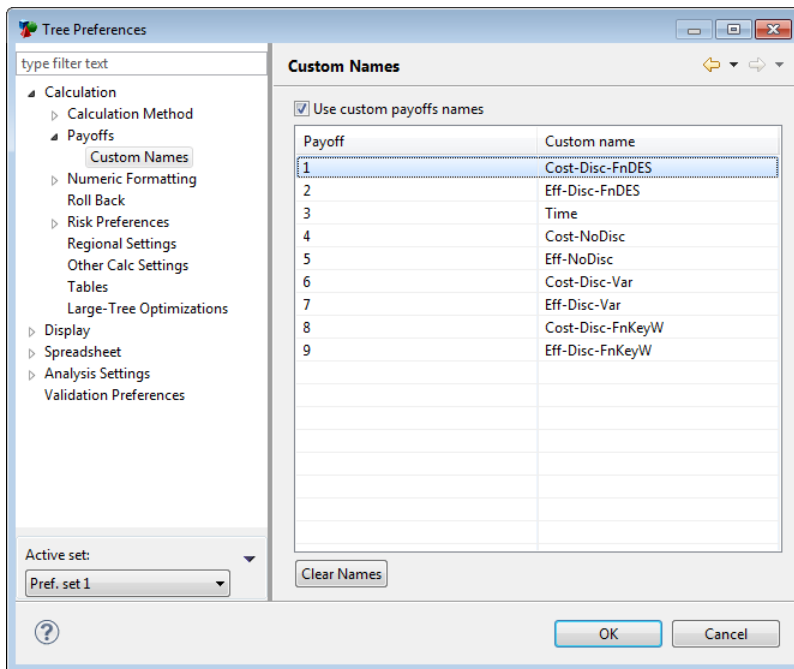
The *Disc* variable is simply the annual discount rate.

The *Disc_i* variable is the instantaneous equivalent of the annual discount rate.

The *Disc_factor_time* variable implements the formula from the figure above. If this is used, it would be placed in a fixed reward because the time is included in the calculation.

The *Disc_factor_rate* divides the *Disc_factor_time* value by *_event_time* to create a discounting adjustment factor that can be applied to rate-based rewards. You might choose to use this variable formula, so you can place the appropriate values in rate-based rewards within the DES Info View. The Osteo DES Model uses this approach in reward sets 1 and 2.

Now, let's consider the time-based rewards in the Osteo DES model with the following reward sets.



DES Model Payoff Names

The figure above shows the time-based rewards at the Process Event node with the three different methods to apply continuous discounting. All three approaches to discounted values will generate the same results.

- Reward sets 1 & 2 use the DiscountContDES function.
- Reward sets 6 & 7 use the variables described above for discounting.
- Reward sets 8 & 9 use the DiscountCont function.

Refer to the Special Features tutorial example model DES.Discounting.trex for a comparison of the Global Discounting function to the manual application of the Discount() function to all rewards. Consider all the payoff sets in the DES Info View to see where global and manual discounting are applied.

38.10 Analysis via Microsimulation

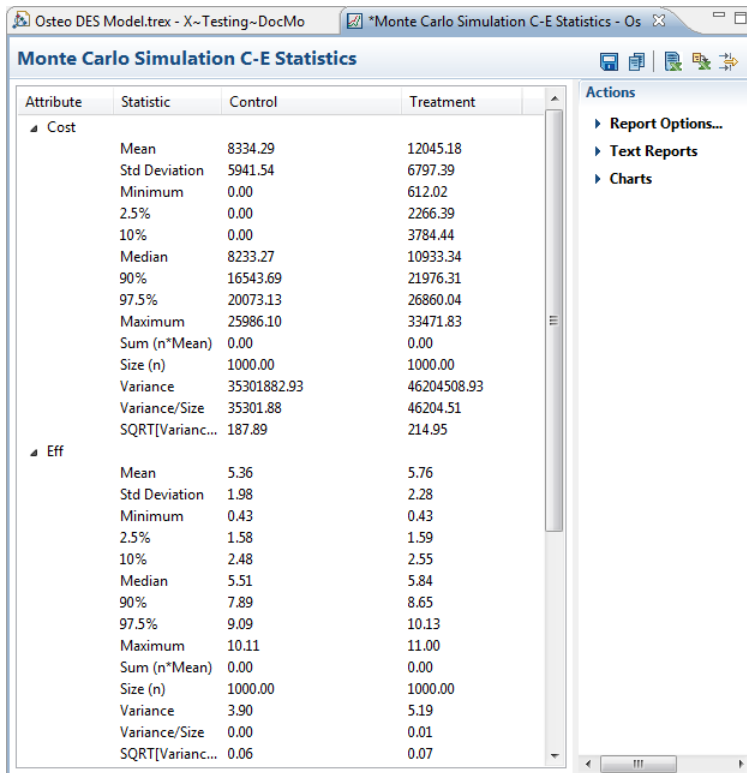
Time-to-event models require individual patients, so they must be analyzed via Microsimulation.

To run Microsimulation on the model:

1. Select the root node.

2. Choose Analysis > Monte Carlo Simulation > Trials (Microsimulation).
3. Choose the number of iterations (1000) and click Begin.

The results are displayed below.



Attribute	Statistic	Control	Treatment
Cost	Mean	8334.29	12045.18
	Std Deviation	5941.54	6797.39
	Minimum	0.00	612.02
	2.5%	0.00	2266.39
	10%	0.00	3784.44
	Median	8233.27	10933.34
	90%	16543.69	21976.31
	97.5%	20073.13	26860.04
	Maximum	25986.10	33471.83
	Sum (n*Mean)	0.00	0.00
	Size (n)	1000.00	1000.00
	Variance	35301882.93	46204508.93
	Variance/Size	35301.88	46204.51
	SQRT(Variance/Size)	187.89	214.95
Eff	Mean	5.36	5.76
	Std Deviation	1.98	2.28
	Minimum	0.43	0.43
	2.5%	1.58	1.59
	10%	2.48	2.55
	Median	5.51	5.84
	90%	7.89	8.65
	97.5%	9.09	10.13
	Maximum	10.11	11.00
	Sum (n*Mean)	0.00	0.00
	Size (n)	1000.00	1000.00
	Variance	3.90	5.19
	Variance/Size	0.00	0.01
	SQRT(Variance/Size)	0.06	0.07

Microsimulation output

The primary output is aggregated data generated from the 1000 individual patient histories. Cost-effectiveness analysis would use the mean values for cost and effectiveness. Many secondary outputs are described in the Individual-Level Simulation and Markov Models Chapter and will not be repeated here.

However, the following is a portion of the text report Statistics - Values, Dists, Trackers, which provides aggregated data on the additional payoffs and trackers.

N_HIP_FRACT_STRATEGY_1	N_HIP_FRACT_STRATEGY_2	N_VERT_FRACT_STRATEGY_1	N_VERT_FRACT_STRATEGY_2
0.72	0.31	2.01	0.90
0.72	0.50	1.42	0.94
0.00	0.00	0.00	0.00
0.00	0.00	0.00	0.00
0.00	0.00	0.00	0.00
1.00	0.00	2.00	1.00
2.00	1.00	4.00	2.00
2.00	1.00	5.00	3.00
3.00	2.00	7.00	5.00
0.00	0.00	0.00	0.00
1000.00	1000.00	1000.00	1000.00
0.52	0.25	2.01	0.88
0.00	0.00	0.00	0.00
0.02	0.02	0.04	0.03

Statistics output

38.11 Calculation trace via the global matrix

Microsimulation output provides the final value for all enabled reward sets and trackers. However, when building/debugging a model, it may be helpful to track each patient's entire path through the model. The Osteo DES Model uses the global matrix and a set of variables to record the patient pathway. Note that these variables are hidden from the model diagram via variable properties to avoid distracting from the main model parameters and formulas. These variables are not added to DES models automatically.



For the calculation trace to function correctly, you must set the tree preferences to run the analysis as a single thread (one CPU). Otherwise, the output data gets written out of order and becomes useless.

Seeding is also helpful when debugging models so that the same results are returned each time.

Both these tree preferences are set for calculation trace in the Osteo DES Model.

38.11.1 Variables used for the calculation trace

All variables within the variable category zTraceOutput are used solely for outputting the global trace.

Variable Name	What it does
_monte_pre_trial_eval	Initializes the counter used to control the row for global matrix updates for each strategy.
_post_analysis	Outputs each strategy's global matrices to a text file after the analysis is complete.
zEventType	Assigns a unique event id for each event node in the model.
zglIndexIncr	Increments the counter used to control the row for global matrix updates for each strategy.

Variable Name	What it does
zgIndexRetr	Retrieves the counter used to control the row for global matrix updates for each strategy.
zgStoreAll	Calls all the individual zgStoreCol... variables described below.
zgStoreCol...	Individual variables to store each piece of critical data in the global matrix for the strategy.
zgStoreMaxTrial	Maximum patient number for the calculation trace. Normally trace info is not needed for every patient when debugging. Running the trace on all patients will significantly slow the analysis. Set to 0 for no trace.

Calculation trace variables

38.11.2 Examining the output

After the analysis is run, two text files are generated, one for each strategy. They can be opened in Excel for orientation by row and column. Output from the *Control* strategy matrix is displayed below (with column headings and shading added).

	A	B	C	D	E	F	G	H	I	J	K	L	M
1	globaln(1)												
2	X:\Testing\DocModels\DES\Osteo DES Model.trex						Control Strategy						
3	Last Distribution Sample												
4	Patient	Event ID	Event Time	Total Time	Cost No Disc	Eff No Disc	Cost Disc	Eff Disc	Hip Frx	Vert Frx	Hip	Vert	Death
5	1	0	3.744567566	3.744567566	0	2.621197296	0	2.459389855	0	0	9.388511241	3.744567566	7.072967153
6	1	2	3.744567566	3.744567566	3000	2.621197296	2637.400629	2.459389855	0	1	9.388511241	1.864873213	7.072967153
7	1	0	1.864873213	5.609440779	3000	3.835229758	2637.400629	3.493171212	0	1	9.388511241	1.864873213	7.072967153
8	1	2	1.864873213	5.609440779	6000	3.835229758	5110.913915	3.493171212	0	2	9.388511241	4.195761059	7.072967153
9	1	0	1.463526374	7.072967153	6000	4.721292531	5110.913915	4.205647727	0	2	9.388511241	4.195761059	7.072967153
10	1	3	1.463526374	7.072967153	6000	4.721292531	5110.913915	4.205647727	0	2	9.388511241	4.195761059	7.072967153
11	2	0	5.337605963	5.337605963	0	3.736324174	0	3.413356533	0	0	6.507465465	5.337605963	15.74197893
12	2	2	5.337605963	5.337605963	3000	3.736324174	2496.752852	3.413356533	0	1	6.507465465	5.736212	15.74197893
13	2	0	1.169859502	6.507465465	3000	4.49790271	2496.752852	4.0345963	0	1	6.507465465	5.736212	15.74197893
14	2	1	1.169859502	6.507465465	10000	4.49790271	8092.708061	4.0345963	1	1	8.303919312	5.736212	15.74197893
15	2	0	4.566352498	11.07381796	10000	7.024693865	8092.708061	5.903902897	1	1	8.303919312	5.736212	15.74197893
16	2	2	4.566352498	11.07381796	13000	7.024693865	10142.33368	5.903902897	1	2	8.303919312	6.80491921	15.74197893
17	2	0	3.737566815	14.81138478	13000	8.94810368	10142.33368	7.13701896	1	2	8.303919312	6.80491921	15.74197893
18	2	1	3.737566815	14.81138478	20000	8.94810368	14347.76719	7.13701896	2	2	7.413117886	6.80491921	15.74197893
19	2	0	0.930594154	15.74197893	20000	9.355167129	14347.76719	7.377699892	2	2	7.413117886	6.80491921	15.74197893
20	2	3	0.930594154	15.74197893	20000	9.355167129	14347.76719	7.377699892	2	2	7.413117886	6.80491921	15.74197893
21	3	0	1.290613143	1.290613143	0	0.9034292	0	0.883667104	0	0	8.468259156	1.290613143	4.829904655
22	3	2	1.290613143	1.290613143	3000	0.9034292	2869.716818	0.883667104	0	1	8.468259156	2.91180546	4.829904655
23	3	0	2.91180546	4.202418603	3000	2.799014554	2869.716818	2.609071874	0	1	8.468259156	2.91180546	4.829904655

DES calculation trace

Note what you can learn about patient 2 from this calculation trace (numbers below are rounded).

- The three time-to-event distributions are sampled prior to the first event.
- The first event is a vertebral fracture at time 5.338 since it is the shortest time-to-event.
- During the time-to-event, time-based utility was accumulated - 3.736 non-discounted utility, 3.413 of discounted utility. There is no time-based cost in the Control strategy.
- The vertebral fracture resulted in a cost of 3,000 (discounted as 2,497).
- The time to vertebral fracture distribution is resampled, drawing a value of 5.736. Therefore, the next vertebral fracture could occur at time $5.338 + 5.736 = 11.074$.

- The second event is a hip fracture at time 6.507. The `_event_time` is $6.507 - 5.338 = 1.170$.
- During the time-to-event, time-based utility was accumulated the same as described above. Discounting will be more significant as this event is further in the future.
- The hip fracture resulted in an additional cost of 7,000 (discounted as 5,596).
- The process continues through another vertebral fracture at time 11.074 and another hip fracture at time 14.811, accumulating the appropriate rate and fixed rewards.
- Finally, the patient dies at time 15.742.

Progression of patient 2 in Control strategy

You may find this type of calculation trace useful as you build/debug your models. A copy of the calculation trace with headings and shading is available in with your tutorial models.

39. State Transition Diagrams

In TreeAge Pro, Markov models are presented and analyzed in decision tree format with health states and transition subtrees. TreeAge Pro also supports state transition diagrams to represent Markov processes, however, analyses can only be run on Markov models in decision tree format.

State transition diagrams can be helpful in providing a simpler representation of the health states and transitions within the model. It can be a good starting point to create a full Markov model and it can also be a good communication tool for an existing Markov model.

This chapter covers the following:

- How to create a state transition diagram
- How to convert a state transition diagram to a Markov model in decision tree format
- How to convert a Markov model in decision tree format to a state transition diagram

39.1 State transition diagrams

State transition diagrams provide a simpler representation of a Markov model than decision trees. While they do not display the level of detail found in a decision tree (i.e., events, probabilities, and rewards), state transition diagrams clearly show the transitions among health states. Even in complex Markov models, where the decision tree is far too large to fit on a single printed page, the corresponding state transition diagram is almost always small enough for simple reproduction and efficient communication.

The design of a state transition diagram is subject to a number of guidelines. Here are the basic ones:

- "*States*" that represent health status.
- "*Arcs*" that represent transitions between any two states.

The combination of states and arcs represent the possible movements of a patient from one health state to another over the entire course of the model timeframe.

39.2 Constructing a state transition diagram

The tutorial in this chapter explains in detail the software commands needed to build a state transition diagram loosely based on the simple three state Markov model described in the Building and Analyzing Markov Models Chapter.

To get started, you will need a new state transition diagram document.

To create a new state transition diagram:

- Choose File > New State Diagram from the menu.

39.2.1 Adding and selecting states

Unlike a new tree, which starts with a root node, a new state transition diagram window is completely blank. The first step in building the diagram is to add the required states. Let's start by creating a *Well* state.

To add a state to a state transition diagram:

- Click on State in the Diagram Editor Palette as seen below.
- Click and hold the mouse within the diagram editor diagram to place the node.
- Drag the mouse down and to the right to resize the new node.



State Transition Diagram Palette

You will see a new, selected state. TreeAge Pro indicates that the state is selected by showing a rectangular outline with resizing markers on every corner and every edge. The node will contain the index value 1 since it is the first state added to the diagram. The node label will also be selected waiting for you to enter the appropriate text.



New unlabeled state selected

The outline around the state can be used to move or resize the state.

When the state is not selected, the rectangular outline will disappear. Since the state label was not entered, it will appear as an ellipsis.



New unlabeled state not selected

To select a state:

- Click inside the state borders.

39.2.2 Entering the state label

You should enter a word or brief phrase in the text box to describe the health state. You can enter the state label when you first create the state, or you can enter/edit the state label later.

To enter/edit the state label:

- Select the state.
- Click on the existing state label or the middle of the state if no text has already been entered.
- Type the state label text (in this case *Well*) in the text area.
- Click outside the state to deselect it.

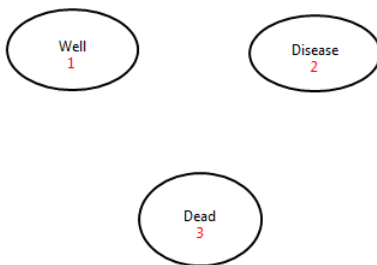


Well state with state label entered



To force a carriage return in a label press *Control + Enter* within the text.

Now add two more states - Disease and Dead.



Added Disease and Dead states

39.2.3 Adding arcs

Arcs are used to show transitions between pairs of states. The arrow head at one end of the arc indicates the direction of the transition.

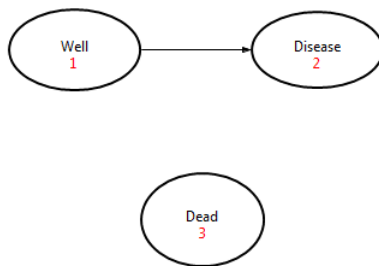
To draw an arc:

- Click on Arc in the State Transition Diagram Editor Palette. See below.
- Click on the *Well* state and drag the arc to the *Disease* state; then release the mouse button.



State Transition Diagram Palette

A new arc will be created pointing from the *Well* state to the *Disease* state.

**Add new arc**

It is frequently useful to add a comment to describe the influence relationship represented by the arc.

To edit an arc label:

- Double-click on the arc.
- Enter text ("Get sick") into the Arc Data dialog Comment field.
- Click OK.

As the arc moves relative to its source/destination states, the label will automatically move with it.

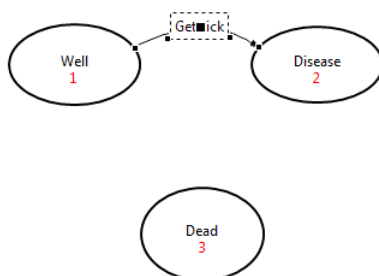
The arc label can also be moved closer/further from the arc.

To move an arc label:

- Click on an arc label and drag it to a new location.

To bend an arc:

- Select the arc.
- Click on the bend point in the middle of the arc and drag it in the direction you wish to bend the arc.
- Release the mouse button.

**Bend an arc**

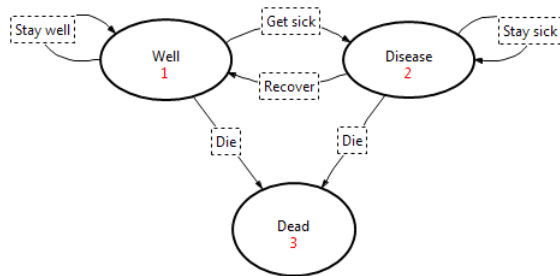
Once an arc has been bent, additional bend points will appear on either side of the bend point that was moved. This allows for more complex arc curvature.

Arcs can be added for transitions from a state back to itself by clicking on the same state for the source and destination of the arc.

Now, add the other required arcs.

- *Recover* arc from *Disease* to *Well*.
- *Die* arc from *Well* to *Dead*.
- *Die* arc from *Disease* to *Dead*.
- *Stay healthy* arc from *Well* to *Well*.
- *Stay sick* arc from *Sick* to *Sick*.

The state transition diagram is now complete.



State Transition Diagram

39.2.4 More editing options

Additional options for editing state transition diagrams are described in this section.

To move a state:

- Select a state and drag its outline (not a resize point) to a new location.

When a state is moved, the arcs attached to it will move accordingly.

To cut a state:

- Select a state.
- Choose Edit > Cut from the menu or press *Control + X* on the keyboard. *Arcs attached to the state will also be cut.*

To resize a state:

- Select a state.
- Click and drag one of the resize marks on the rectangular outline around the state.

To select multiple states:

- Select one state then hold down the Shift key and select another state.
- ... Or...

- Click on open space in the diagram and drag to create a rectangle that surrounds all elements you wish to select

To delete an arc:

- Select the arc.
- Press the *Delete* button on the keyboard.

To bend an arc:

- Click on the arc points and drag it to another location. Rather than moving the entire arc, it will be split into two line segments, each terminated at the new drag location.
- You can bend this further by dividing up those line segments using the same technique.

To arrange all states evenly:

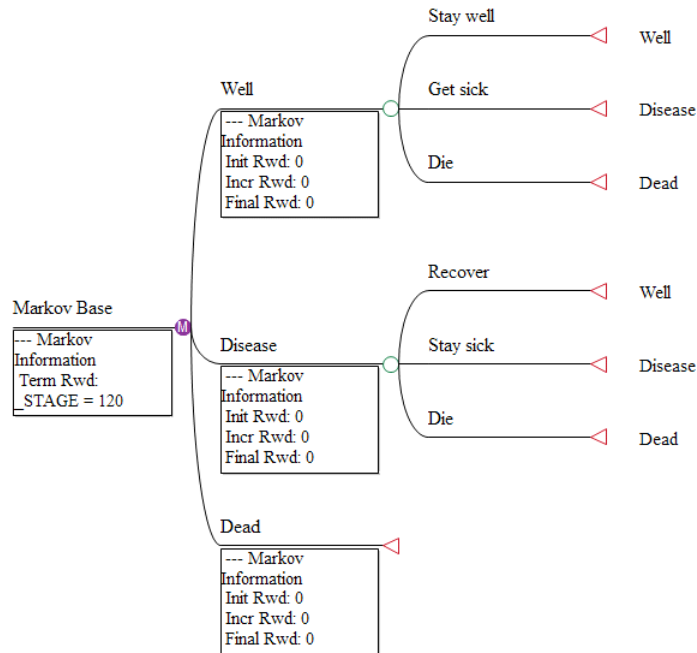
- Choose Edit > Auto-arrange states.

39.3 Converting a state transition diagram to a decision tree

State transition diagrams cannot provide all the details required for most Markov models. However, you can create a state transition diagram to conceptualize the Markov model and then convert it to the decision tree format.

To convert the diagram:

- Open and activate the state transition diagram editor window.
- Choose Edit > Convert to tree from the menu.



State transition diagram converted to a markov model

The resulting Markov model includes all the states from the state transition diagram. It also includes all the transitions emanating from each state and the arc labels become the node labels for the transitions.

Markov rewards and transition probabilities could then be added to the model. The transition subtrees may also be expanded to add events.

The Markov model can be incorporated into a larger decision tree via copy/paste subtree.

39.4 Converting a Markov model to a state transition diagram

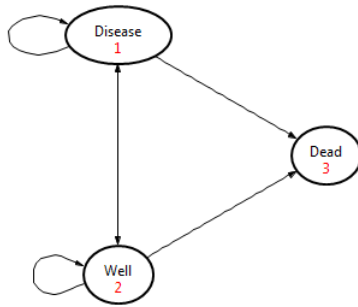
Since analyses are run on the Markov model in decision tree format, you might ask why you would want to convert a Markov model to a state transition diagram. Well, you might want a simpler representation of the Markov model for communication purposes. With a state transition diagram, it's easier to see the flow of patients among the health states.

To illustrate this process, you can open the Health Care tutorial example model "Three-State Markov."

To convert a Markov model to a state transition diagram:

- Right-click on a Markov node in the decision tree.
- Choose Convert to STD from the context menu.

The result is a new state transition diagram with all states and transition arcs. Note that there may be many paths from one state to another, but only one arc will be created.



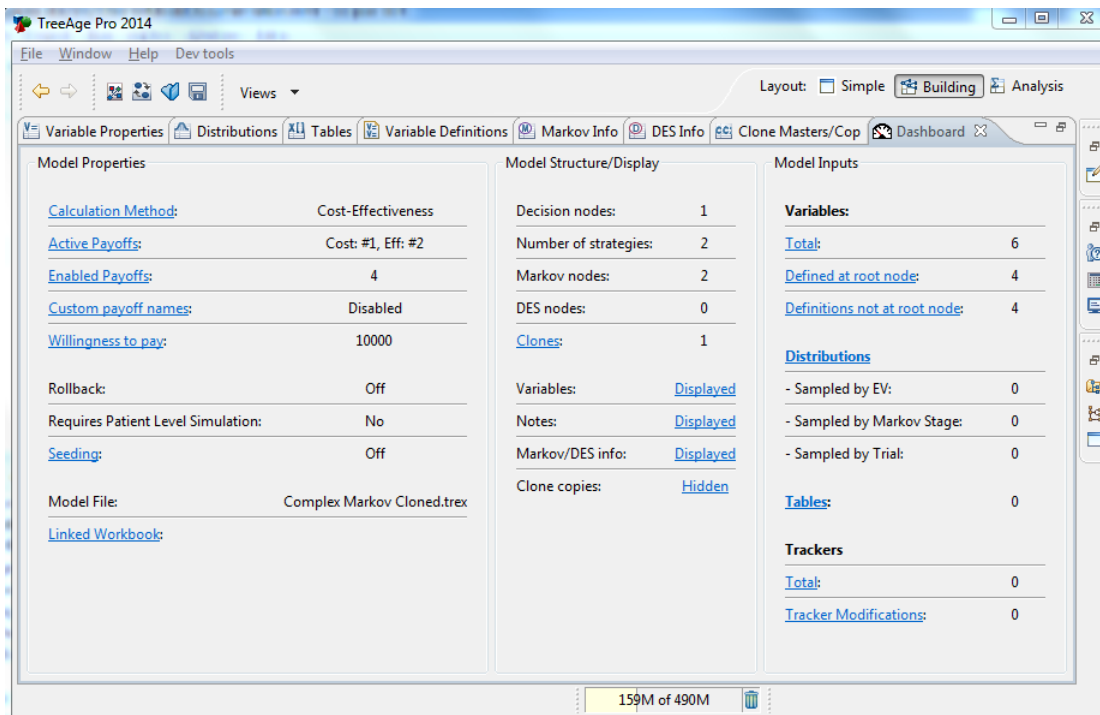
Markov model converted to state transition diagram

40. Model Dashboard and Documentation

This chapter describes the Dashboard View which gives an overview of your model and Model Documentation which is a mechanism used to document your model.

40.1 Using the Model Dashboard

TreeAge Pro provides a View to give you an overview of your model. You access the Dashboard through the Views menu. The figure below shows what the Dashboard looks like.

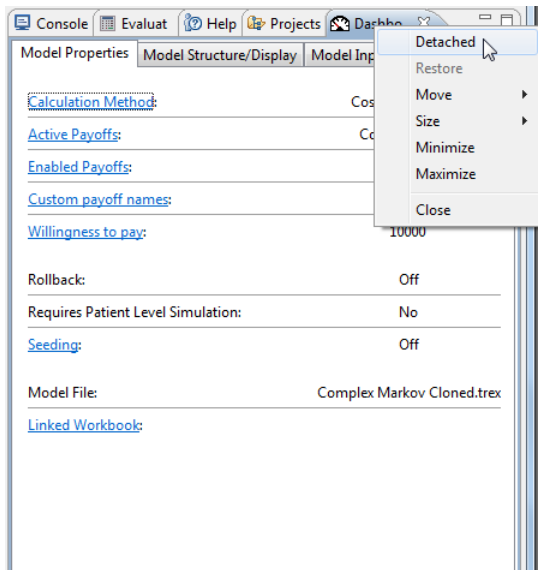


Dashboard View - maximised to fill the whole screen

The Dashboard is split into 3 sections, each providing different information about your model. Selecting any of the text highlighted in blue will take you to where you can edit the details or it will toggle between settings (such as Displayed/Hidden). The 3 sections are:

- *Model Properties*: Access all the main properties of the model including Calculation Method, Payoffs, willingness to pay and any linked Workbooks.
- *Model Structure/Display*: Details about the number of strategies, number and type of nodes, and display settings such as whether to show variables, notes, Markov info and clone copies is found here.
- *Model Inputs*: Any information entered into the model can be updated through this pane including variables, distributions, tables and trackers. Selecting any of the blue text will take you to the appropriate View.

When the Dashboard view is not full size, each of the sections appear as selectable tabs to allow access to all the information by clicking through each tab, as in the figure below. This pane can also be detached from TreeAge Pro, right-clicking on the tab and selecting "detach".



Dashboard View with tabs for each of the 3 panes. Right-click and select Detached to detach and move independently of the main windows.

40.2 Create Model Documentation Help File

TreeAge Pro provides Help html files to describe the software and its functions. You can create a similar Help html file to describe your model.

There are three steps to this process.

- Create an Excel workbook for model documentation based on the model structure.
- Edit the Excel workbook to add documentation text.
- Import the Excel workbook to create a Help html file.

Creating model documentation

We will use the Get Started tutorial example model Three Vars to demonstrate this technique.

To create an Excel workbook for model documentation:

- Open the model.
- Choose File > Model Help/Documentation > Create Help Workbook > Simple Model Docs Template.
- Select the model input types that exist in your model and what you wish to document.



Different Help Workbooks are available for cost-effectiveness and CE Markov models.

This creates a workbook file next to the model file within the same folder with the same name, but with "_ModelDocs.xls" added to the end. The workbook contains placeholder cells for entering model documentation.

To open the workbook for editing in Excel:

- Select the applicable model in the Tree Diagram Editor.
- Choose File > Model Help/Documentation > Open Documentation Spreadsheet.

The cells with labels and "<title>" type tags should be left alone. The cells to the right of these "label" cells are used for entry of descriptive text. See below.

	A	B
1		
2		
3		
4		
5	<section>I. Structure	
6		
7	<subsection>A. Strategies	
8		
9	<question>1. What is the important background for the decision in this model?	Comparing two investment strategies.
10		
11	<question>2. What strategies are chosen for comparison?	Risky investment with unknown return. CD with fixed return.
12		
13	<question>3. Describe the reasons for the inclusion/exclusion of strategies:	No other strategies were used for this simple example.
14		
15		
16	<subsection>B. Events:	
17		
18	<question>4. List the key events in the model. Are they different for different strategies? If so, why?	The risky strategy is dependent on market forces which could result in any of three
19		
20		
21	<subsection>C. Other assumptions:	
22		
23	<question>5. What is the time frame of the model (months, years, etc.)?	One year horizon.
24		
25		
26	<subsection>C. Other information:	
27		
28	<question>6. Is the structure of this model based on other, similar, published or otherwise accessible models?	This is just an example model. No relation to published results.
29		
30	<question>7. What sources (experts, panels, studies or publications) were used to determine the structure of the model?	No sources were used.
31		
32		
33		
34		
35		
36		
37		
38		

Model Documentation Workbook

Once the text in the workbook has been updated, it can be imported back into TreeAge Pro to generate the Help html file.

To create the Help file:

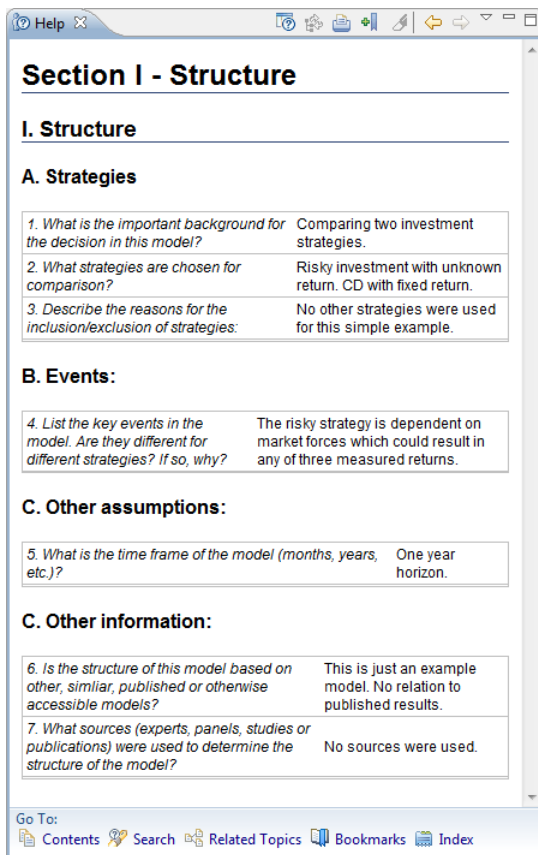
- Select the applicable model in the Tree Diagram Editor.
- Choose File > Model Help/Documentation > Export Workbook to HTML Help

This creates an HTML file next to the model file within the same folder with the same name, but with "_ModelDocs.html". This Help file is then available for documentation of the model.

To open the Help file:

- Select the applicable model in the Tree Diagram Editor.
- Press the *F1* key.

The Help file will look like this.



Model Documentation Help

41. Model Validation

This chapter describes a mechanism used to validate your model by looking for commonly found flaws within the model.

41.1 How model validation works

TreeAge Software has identified a set of common flaws in models; the model validation process checks for these flaws in your model and presents any occurrences of these flaws back to you.

The specific validation checks are:

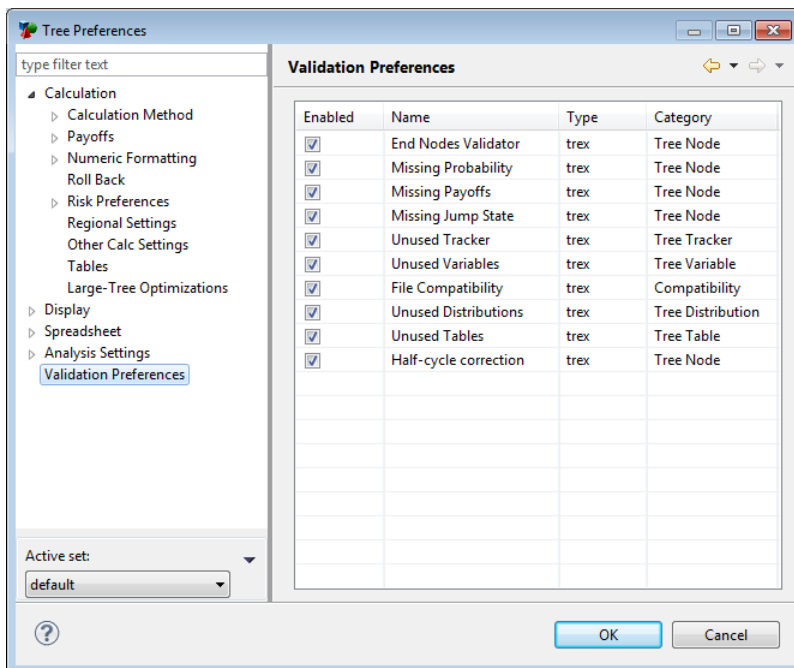
- *End Nodes Validator*: find end nodes that are not terminal nodes.
- *Missing Probability*: find branches of chance nodes with no probability expression.
- *Missing Payoffs*: find terminal nodes (non-Markov) that are missing payoffs.
- *Missing Jump State*: find terminal nodes (Markov) that are missing a jump state.
- *Unused Tracker*: find trackers that are not used in the model.
- *Unused Variables*: find variables that are not used in the model.
- *Unused Distributions*: find distributions that are not used in the model.
- *Unused Tables*: find tables that are not used in the model.
- *File compatibility*: identify features that are not supported by older versions of TreeAge Pro.

Model Validation Checks

More validation checks will be added to TreeAge Pro in future releases.

41.2 Determine which validators to run

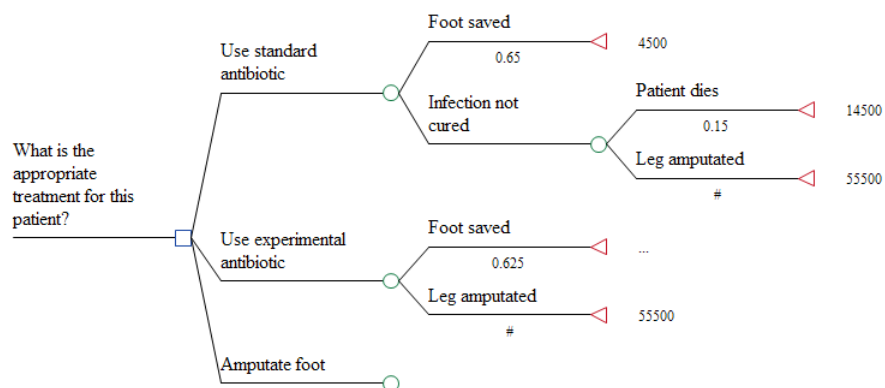
Tree Preferences provide the ability to turn on/off specific validator checks.



Uncheck any of the model validators to skip that check in the validation process.

41.3 Run the model validation

To illustrate the model validation process, we will open Special Features tutorial example model ModelValidationTest. As you can see below, it has a few model flaws: missing probabilities, missing payoffs and non-terminal end nodes.

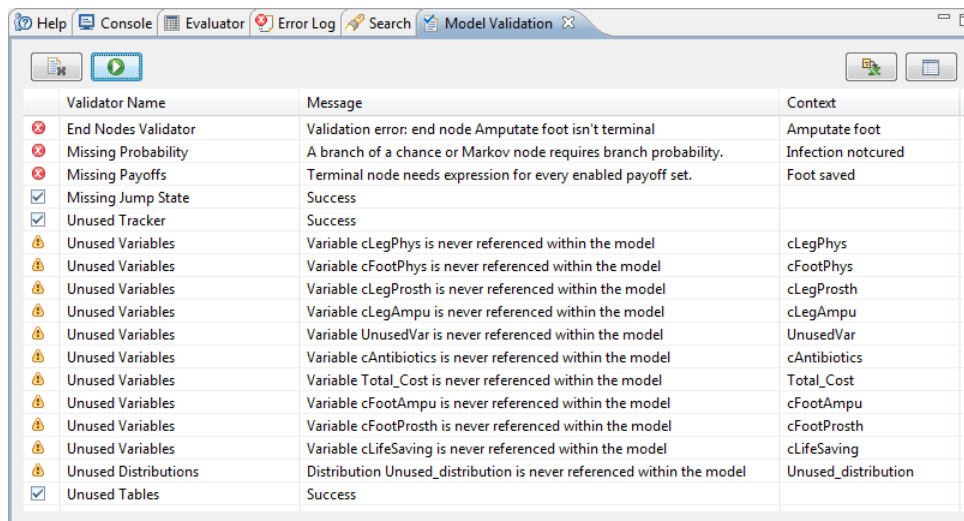


ModelValidationTest model

To run the model validation:

- Open/activate the model in the Tree Diagram Editor.
- Open the Model Validation View from the Views list in the toolbar.
- Click the Validate button (green circle with white triangle).

The Model Validation View will display the results from all the validation checks. See below.



Validator Name	Message	Context
End Nodes Validator	Validation error: end node Amputate foot isn't terminal	Amputate foot
Missing Probability	A branch of a chance or Markov node requires branch probability.	Infection notcured
Missing Payoffs	Terminal node needs expression for every enabled payoff set.	Foot saved
Missing Jump State	Success	
Unused Tracker	Success	
Unused Variables	Variable cLegPhys is never referenced within the model	cLegPhys
Unused Variables	Variable cFootPhys is never referenced within the model	cFootPhys
Unused Variables	Variable cLegProsth is never referenced within the model	cLegProsth
Unused Variables	Variable cLegAmpu is never referenced within the model	cLegAmpu
Unused Variables	Variable UnusedVar is never referenced within the model	UnusedVar
Unused Variables	Variable cAntibiotics is never referenced within the model	cAntibiotics
Unused Variables	Variable Total_Cost is never referenced within the model	Total_Cost
Unused Variables	Variable cFootAmpu is never referenced within the model	cFootAmpu
Unused Variables	Variable cFootProsth is never referenced within the model	cFootProsth
Unused Variables	Variable cLifeSaving is never referenced within the model	cLifeSaving
Unused Distributions	Distribution Unused_distribution is never referenced within the model	Unused_distribution
Unused Tables	Success	

Model Validation View

The results are presented in a grid with a row for each validation check/problem.

The first column indicates whether the row represents an error, warning or a successful check. A successful check means that no occurrences of problems were found.

The second column displays the validator name.

The third column displays a message related to the problem.

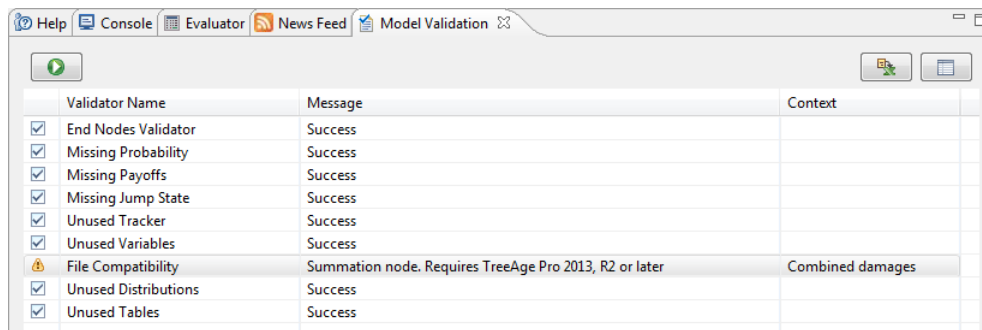
The fourth column displays the context for that error/warning. The context is used to provide a node label or model input name associated with the problem.

Double-click on any error/warning line in the grid to move either to the tree diagram editor to the highlighted node with the problem or to the proper model input view for the model input with the problem. This gives you immediate access to the context where the problem can be fixed.

41.4 File compatibility

Model validation identifies features that are used, but are not supported by earlier software versions. If you have used any such features, you will need to ensure that your colleagues have software that supports those features or they will not be able to open the model. Alternatively, you can modify your model to remove those features.

The following model validation output is from the Legal tutorial example model LegalTreeSummation.



Validator Name	Message	Context
<input checked="" type="checkbox"/> End Nodes Validator	Success	
<input checked="" type="checkbox"/> Missing Probability	Success	
<input checked="" type="checkbox"/> Missing Payoffs	Success	
<input checked="" type="checkbox"/> Missing Jump State	Success	
<input checked="" type="checkbox"/> Unused Tracker	Success	
<input checked="" type="checkbox"/> Unused Variables	Success	
<input checked="" type="checkbox"/> File Compatibility	Summation node. Requires TreeAge Pro 2013, R2 or later	Combined damages
<input checked="" type="checkbox"/> Unused Distributions	Success	
<input checked="" type="checkbox"/> Unused Tables	Success	

Model validation - file compatibility

As indicated in the validation output above, the summation node feature was introduced with TreeAge Pro 2013, R2.

42. Preferences

This chapter describes TreeAge Pro preferences - Tree Preferences and Application Preferences.

The first section of this chapter describes Tree Preferences, which are specific to the active tree model, and are stored with the tree document. The last few sections describe Application Preferences, which are associated with the TreeAge Pro application and apply generally to the functioning of the software and to all models.

42.1 Tree Preferences

Tree Preferences provide control over many settings and options for the active model, including how it is calculated and how it is displayed. Although they are not part of the model structure or numerical data, Tree Preferences are important elements of a model which are stored within the model document.

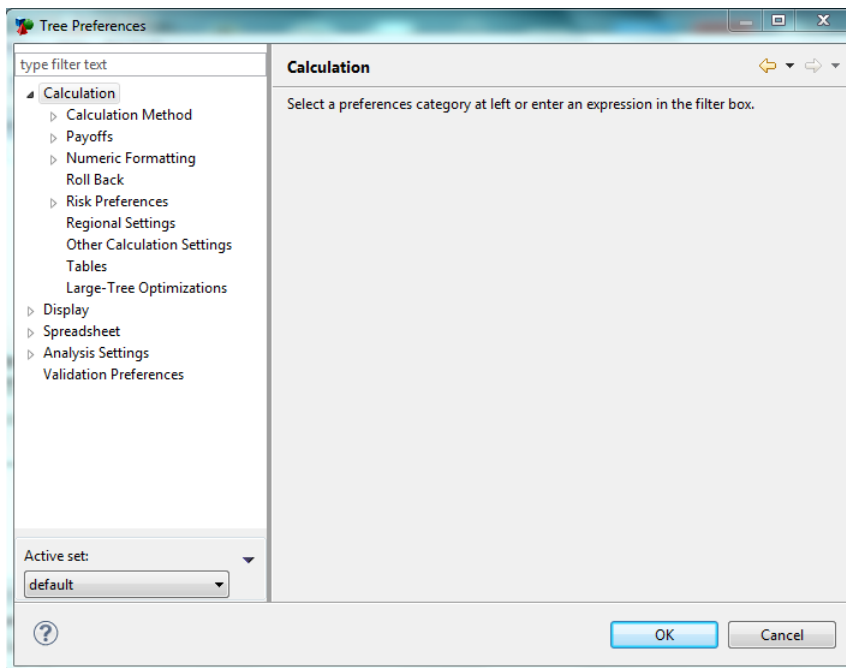
Tree Preferences can be broken into a few main categories:

- Calculation preferences
- Display preferences
- Spreadsheet preferences
- Analysis settings

Some frequently used Tree Preferences can be set by using the Configure Model wizard. The wizard prompts the user through specific elements of the Tree Preferences as listed below:

- Calculation method
- Additional options associated with the selected calculation method
- Payoffs
- Discounting
- Custom payoff names
- Numeric formatting

It is also possible to edit any Tree Preferences directly using the Tree Preferences Dialog.

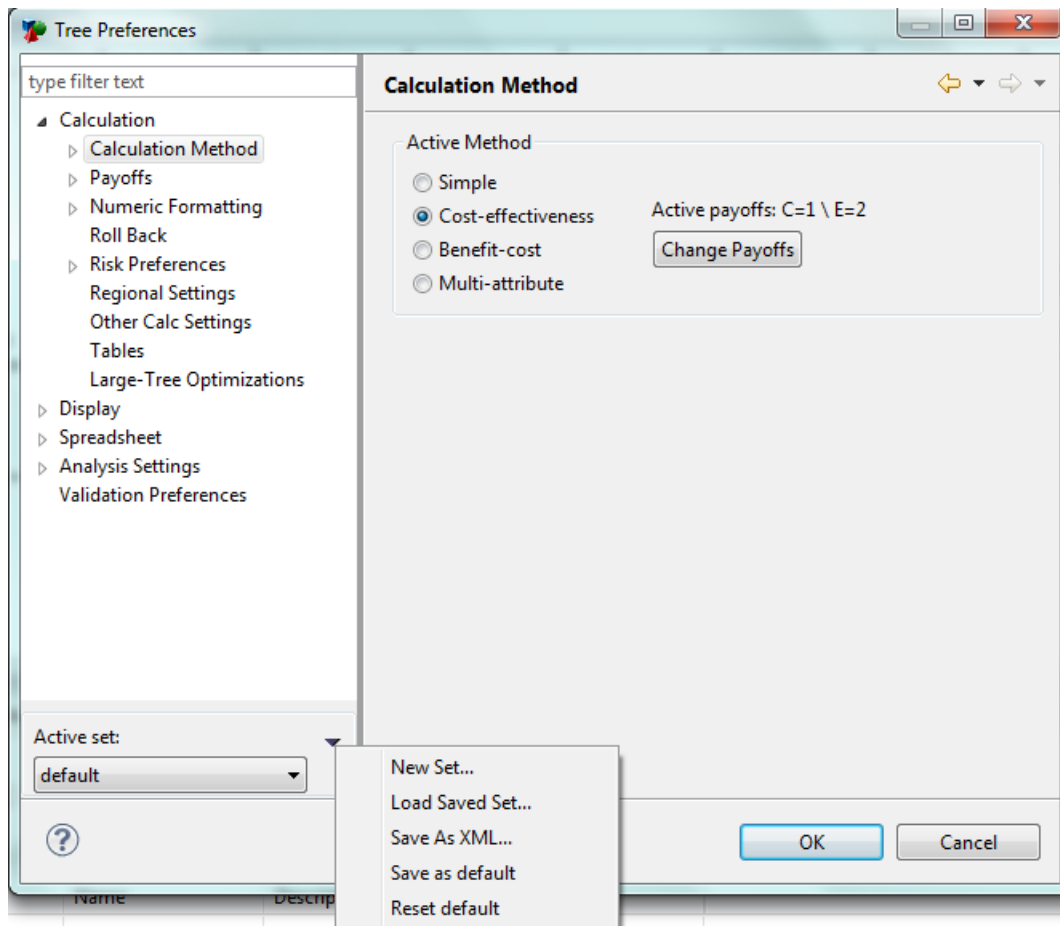


Tree Preferences dialog (Windows)

42.2 Tree Preference Sets

TreeAge Pro allows you to save and load Tree Preference Sets, which include all the preference settings from the Tree Preferences Dialog. For example, you might regularly create two kinds of trees, cost-effectiveness trees and simple trees. You could save a Preference Set for each of these types from a model that you have already created. Then you could load the appropriate Preference Set when creating a new tree.

The controls for Preference Sets are at the bottom left corner of the Tree Preferences Dialog.



Tree Preference Sets with controls listed

In the figure above, the down arrow control has been clicked to show the Tree Preference Set controls, which are described below:

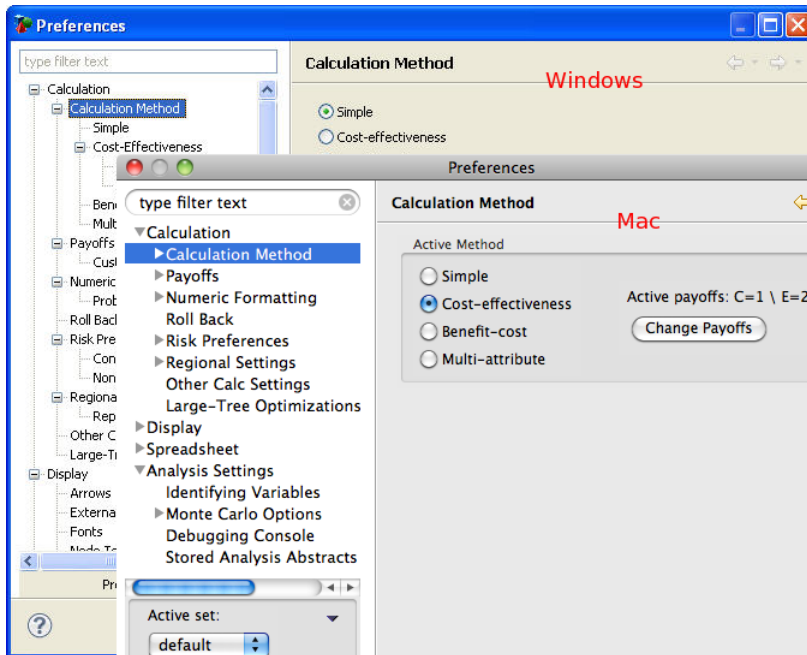
- *New Set*: Create a new preference set for the selected model. The new preference set can be copied from an existing preference set already saved with the model.
- *Load Saved Set*: Load a set of preferences from an XML file into a preference set for the selected model. You will be asked to provide a name for the preference set within the context of the selected model.
- *Save AS XML*: Save the current preference set as an XML file. It can then be loaded into other models.
- *Save as default*: Save the current preference set as the default for new models.
- *Reset default*: Reset the default preference set for new models to the set that comes with the original software installation.

Tree Preference Set Controls

42.3 Tree Preferences Dialog

The Tree Preferences Dialog is used to edit Tree Preferences for the active model. You can open the Tree Preferences Dialog by first selecting a model and then using one of the following methods:

- Choose Tree > Tree Preferences from the menu.
- Click the "preferences" icon on the toolbar.
- Windows/Linux: Pressing F11 on the keyboard. Mac OS: Set up a keyboard shortcut, if desired, in the Application Preferences



Tree Preferences Dialog - Mac + Windows

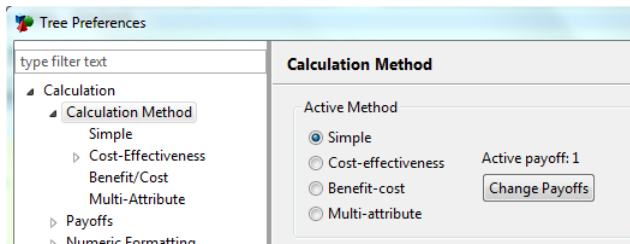
42.3.1 Tree Preference Categories

Tree Preferences are broken up into a number of categories as shown on the left side of the dialog. The list of available categories changes depending on what, if any, type of document is currently active. The list of categories can be filtered using the filter text input. Choosing a category from the list changes the page of options that appear in the right side of the dialog. The options in each category are described in this chapter.

42.4 Calculation Method

This is the first category related to *calculation* preferences.

A model's Calculation Method controls the primary manner in which the model is calculated. TreeAge Pro supports four Calculation Methods.



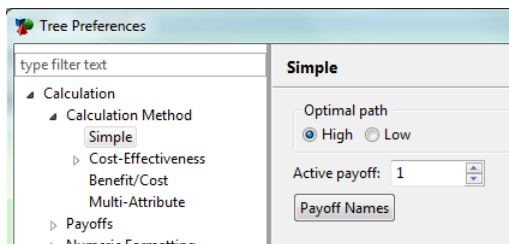
Calculation Method Selection

- Simple: Calculate values using a single payoff set.
- Cost-effectiveness: Calculate separate cost and effectiveness values using two selected payoff sets (by default, #1 and #2). Also enables incremental cost-effectiveness calculations in a variety of analyses, including sensitivity analysis and Monte Carlo simulation.
- Benefit-cost: Calculate values based on a cost payoff set subtracted from a benefit payoff set.
- Multi-attribute: Calculate values based on a weighted combination of several payoff sets.

Calculation Methods

42.4.1 Calculation Method - Simple

There are a few options available to control the Simple Calculation Method.

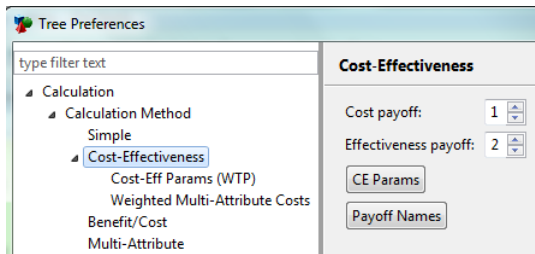


Tree Preferences - Simple Calculation Method

- *Optimal path*: Select *High* when your tree is to be calculated on the basis of profit, utility, cash flow, quality of life or other attributes that should be maximized. Select *Low* when payoffs are costs or other attributes that should be minimized.
- *Active payoff*: You can assign up to nine payoffs at each terminal node. Changing the "Active payoff" determines which of these payoffs are to be used in calculating the model.

42.4.2 Calculation Method - Cost-Effectiveness

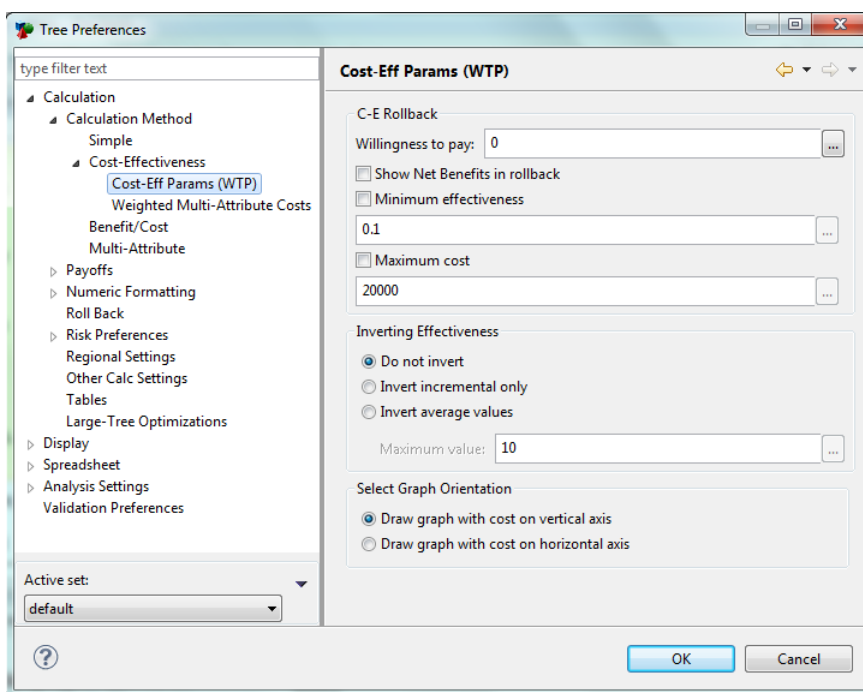
There are a few options available to control the Cost-Effectiveness Calculation Method. There are also two subcategories: Cost-Effectiveness Parameters and Weighted Multi-Attribute Costs.



Tree Preferences - Cost-Effectiveness Calculation Method

- *Cost payoff*: Select the payoff to use for cost calculations within the model.
- *Effectiveness payoff*: Select the payoff to use for effectiveness calculations within the model.

Cost-Effectiveness Parameters (WTP)

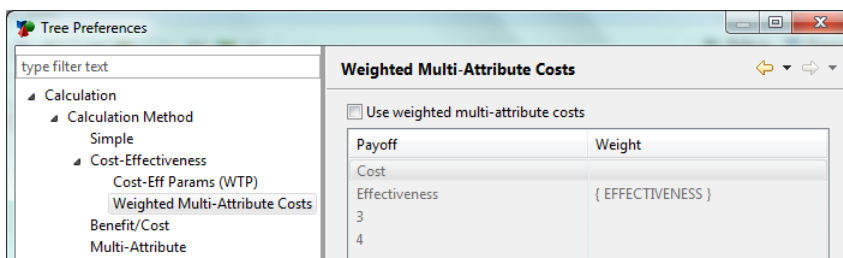


Tree Preferences - Cost-Effectiveness Calculation Method - C/E Parameters

- *Willingness to pay*: Enter the weighting to place on effectiveness when balancing against cost.
- *Minimum effectiveness*: Check this box to activate the minimum effectiveness value in CE calculations.
- *Minimum effectiveness value*: Enter the minimum effectiveness value. Strategies that do not meet this minimum effectiveness requirement are eliminated from CE calculations.
- *Maximum cost*: Check this box to activate the maximum cost value in CE calculations.
- *Maximum value*: Enter the maximum cost value. Strategies that do not meet this maximum cost requirement are eliminated from CE calculations.
- *Inverting effectiveness*: The inversion options allow you to invert effectiveness when the measurement of effectiveness needs to be minimized (e.g., cases avoided) rather than maximized (e.g., life years).

- *Do not invert*: Do not invert the effectiveness value. This is the default value which assumes that you want to maximize the effectiveness measure.
- *Invert incremental only*: Invert only the incremental values among strategies. This is usually the preferred method to handle minimizing effectiveness values
- *Invert average values*: Invert average cost-effectiveness values. This inverts all effectiveness calculations. When selected, you must enter a maximum effectiveness value.
- *Maximum value*: Enter a fixed maximum effectiveness value from which to subtract all nodes' calculated effectiveness values during cost-effectiveness calculations.
- *Select Graph Orientation*: Choose which graph format you wish to output.
- *Draw graph with cost on vertical axis*: Places cost on the Y-axis of the graph. This is the default setting.
- *Draw graph with cost on horizontal axis*: Places cost on the X-axis of the graph.

Weighted Multi-Attribute Costs

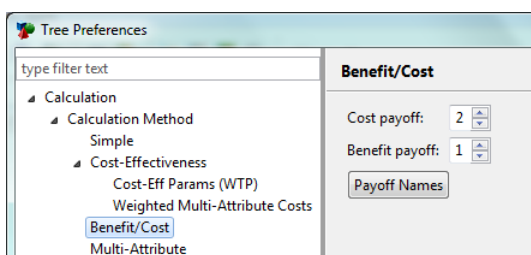


Tree Preferences - Cost-Effectiveness Calculation Method - Weighted Multi-Attribute Costs

- *Use weighted multi-attribute costs*: Check this box to activate weighted multi-attribute costs.
- *Payoff/weight grid*: Enter a weight for each payoff set.

42.4.3 Calculation Method - Benefit/Cost

There are a few options available to control the Benefit/Cost Calculation Method.

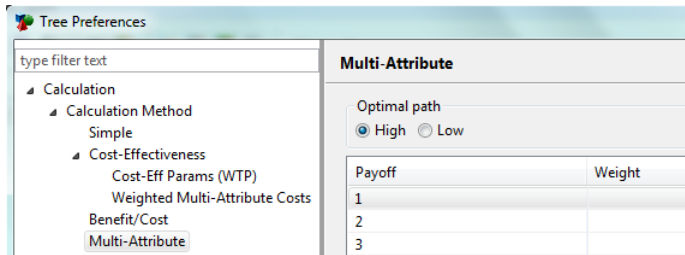


Tree Preferences - Benefit/Cost Calculation Method

- *Cost payoff*: Select the payoff to use for cost calculations within the model.
- *Benefit payoff*: Select the payoff to use for benefit calculations within the model.

42.4.4 Calculation Method - Multi-Attribute

There are a few options available to control the Multi-Attribute Calculation Method.

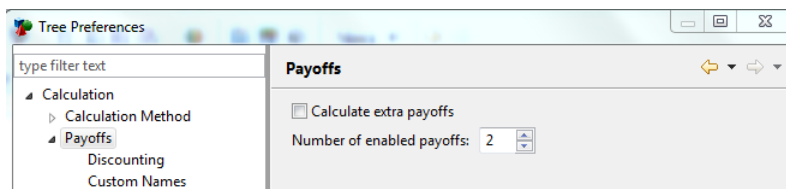


Tree Preferences - Multi-Attribute Calculation Method

- *Optimal path*: Select High when the weighted multi-attribute calculated values should be maximized. Select Low when they should be minimized.
- *Payoff/weight grid*: Enter a weight for each payoff set.

42.5 Payoff Preferences

The Payoffs Tree Preferences category allows you to control the payoffs that are enabled in the model and how the model discounts rewards for Markov and DES models. This preference also controls the payoffs available within other Tree Preferences.

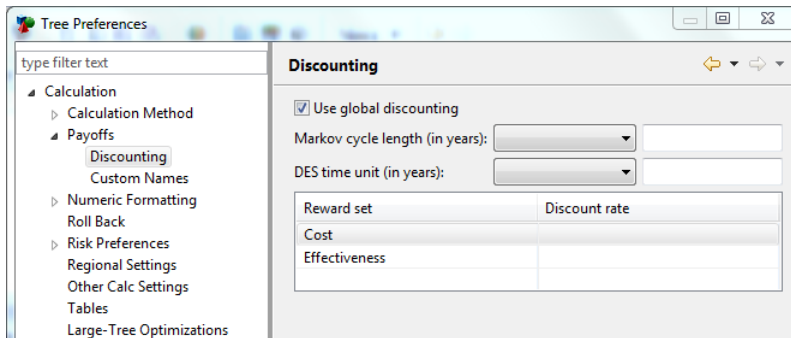


Tree Preferences - Payoffs

- *Calculate extra payoffs*: Check this box to calculate additional payoffs in addition to the ones specified as active within the model's calculation method. Additional rewards are calculated in Markov Cohort Analysis and Monte Carlo Simulation.
- *Number of enabled payoffs*: Select the number of payoffs to enable, starting with payoff 1.

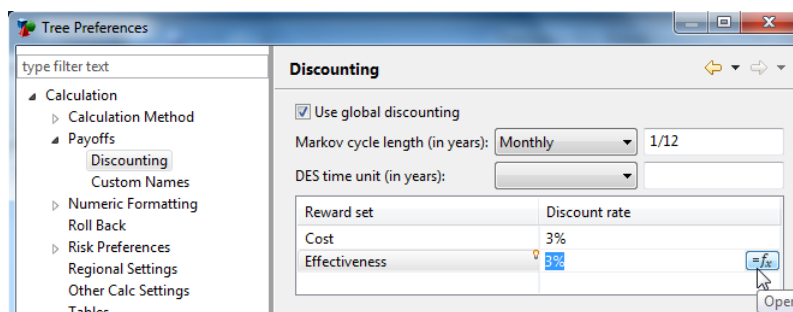
42.5.1 Discounting Rewards

The Payoffs - Discounting Tree Preferences category allows you to discount entire rewards sets for Markov and DES models. The Reward set is populated based on the number of enabled payoffs. For more information about the formula used in discounting, review the appropriate section about Markov models or DES models.



Tree Preferences - Discounting

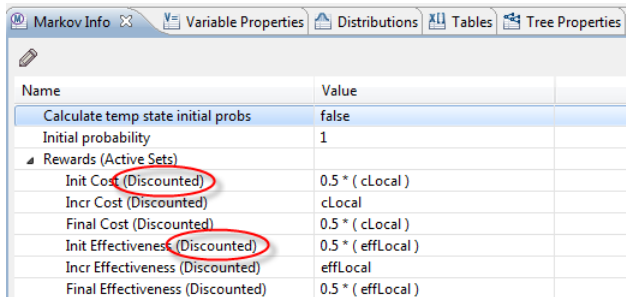
- *Use global discounting*: Check this box to enable Global Discounting.
- *Markov cycle length (in years)*: There are two options to set the time interval used for discounting.
 1. Select from the dropdown menu one of the most commonly used Markov cycle lengths appropriate to your model (Annually, Quarterly, Monthly, Daily, Weekly or Hourly). This pre-fills the time interval used for discounting in Annual units.or....
 2. Manually enter the time interval to be used for discounting in Annual units into the blank field. For example, 4 monthly cycles would be entered as 1/3.
- *DES time unit (in years)*: This refers to the units returned by time-to-event distributions. The two options to set the time interval for DES are the same as for Markov cycle length as described above.
- *Reward set/Discount rate grid*: This is where you set the discount rate to be applied to each reward set. The reward sets are generated from the enabled payoffs. If the model calls for a discount rate to only be applied to one reward set (unlikely) do not enter a discount rate for that reward set.



Enabling global discounting and apply to Markov model with Monthly cycle lengths and 3% discount rate.

- Note the ability to either enter a numeric value (percentage) to describe the Discount Rate or to enter a more complicated formula based on model variables using the "=fx" button.
- The figure below shows a model's Markov Info View highlighting those rewards which have been discounted with global discounting. Note there is no change to the reward in the value

column, but the word *Discounted* next to the reward name indicates Global Discounting has been applied.



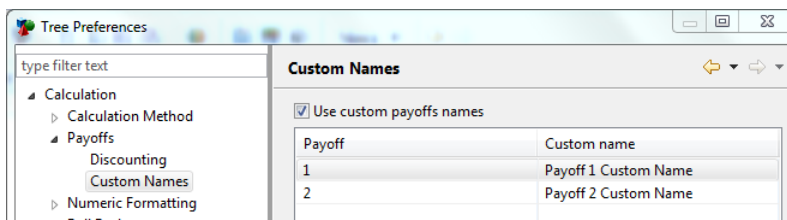
Name	Value
Calculate temp state initial probs	false
Initial probability	1
▲ Rewards (Active Sets)	
Init Cost (Discounted)	$0.5 * (cLocal)$
Incr Cost (Discounted)	cLocal
Final Cost (Discounted)	$0.5 * (cLocal)$
Init Effectiveness (Discounted)	$0.5 * (effLocal)$
Incr Effectiveness (Discounted)	effLocal
Final Effectiveness (Discounted)	$0.5 * (effLocal)$

Markov Info view illustrating where rewards have been discounted.

- *DES time unit (in years)*: There are two options to set the time interval used for discounting. This time interval should be consistent with the units used in the distributions for the time to event sampling.
 1. Select from the dropdown menu one of the most commonly used DES time to event distribution time units appropriate to your model (Annual, Quarterly, Monthly, Daily, Weekly or Hourly). This pre-fills the time interval used for discounting in Annual unitsor....
 2. Manually enter the time interval to be used for discounting in Annual units into the blank field. For example, 4 monthly cycles would be entered as 1/3.

42.5.2 Custom Payoff Names

The Payoffs - Custom Names Tree Preferences category allows you to create custom names for the enabled payoffs.



Tree Preferences - Custom Payoffs Names

- *Use custom payoff names*: Check this box to enable custom payoff names.
- *Payoff/Custom name grid*: Enter a custom name for each payoff set.

You can enter custom payoff names for all enabled payoffs. When entering payoffs at Terminal Nodes, the custom payoff names will be displayed in place of the payoff set index.

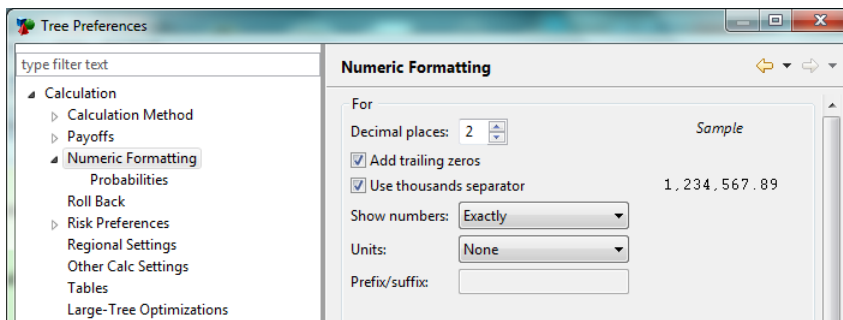
42.6 Numeric Formatting

The Numeric Formatting Tree Preferences category allows you to control how numerical output is displayed within TreeAge Pro.

You can enter separate numeric formatting settings for each payoff, but only active payoff(s) are presented. As the Calculation Method preferences change, so too will the active payoff(s) or combined payoff output presented under Numeric Formatting change. In the figure below, the Calculation Method is Simple and payoff 1 is active. Therefore, Numeric Formatting options are only presented for payoff 1.

If the Calculation Method were Cost-effectiveness, then the active cost and the active effectiveness payoffs would be displayed, as well as numeric formatting for cost-effectiveness (a combination of the two payoffs).

If your model calculates additional payoffs beyond the active payoff(s), you can temporarily change the active payoff to allow you to enter numeric formatting for that payoff.



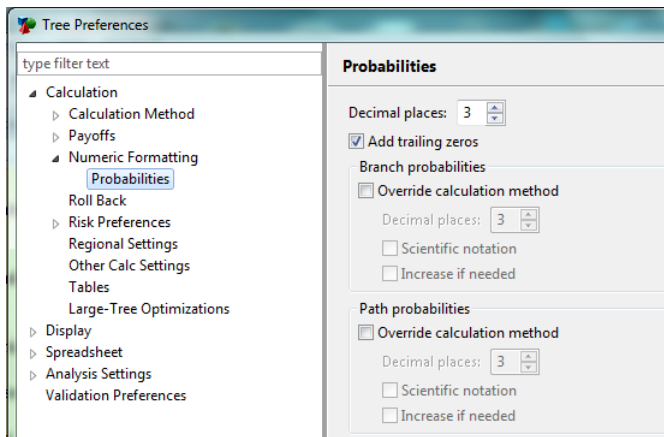
Tree Preferences - Numeric Formatting

- *For payoff _*: Indicates the payoff for which these Numeric Formatting preferences apply.
- *Decimal places*: Select the number of decimal places to display for the selected payoff.
- *Use thousands separator*: Check this box to separate thousands using the defined separator for the model or for the computer.
- *Show numbers in*: Select what order of magnitude you want to show numbers. Options include *exactly*, *in thousands*, *in millions*, *in billions* and *in percent*. The option you choose will depend on the size of the numbers used in the model.
- *Units*: Select the units that apply to that payoff.
None: Show calculated values with no units.
Currency: Show calculated values as currency. Note that the currency normally derives from the standard operating system settings on the computer. However, the currency can be overridden within the Regional Settings Tree Preferences.
Custom prefix: Display custom text before the calculated values.
Custom suffix: Display custom text after the calculated values.
- *Prefix/suffix*: Enter the custom text to be used when the *Units* are defined as *Custom prefix* or *Custom suffix*.

- *Sample*: Shows how calculated values will be displayed based on the Numeric Formatting preferences.

42.6.1 Probabilities

The Numeric Formatting Tree Preferences for Probabilities allows you to control how the probability numerical output is displayed within TreeAge Pro.

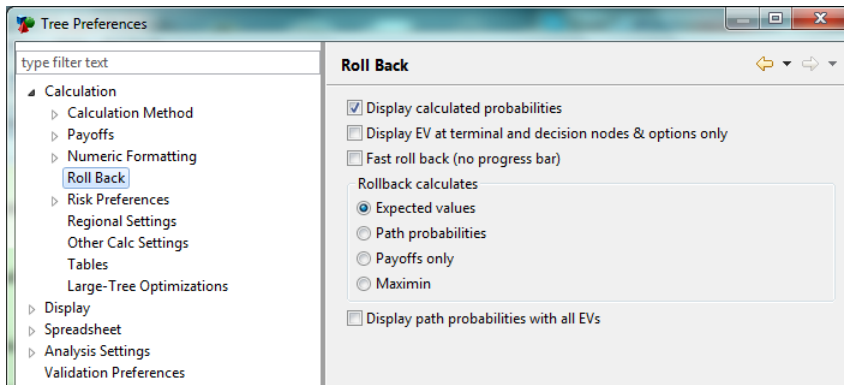


Tree Preferences - Numeric Formatting, Probabilities

- *Decimal places*: Select the number of decimal places to display for all probabilities.
- *Branch probabilities*: Check the "Override calculation method" box and select the number of decimal places to use a different probability format for branch probabilities. Checkboxes further specify whether to use scientific notation and/or increase the number of decimal places as needed.
- *Path probabilities*: These are the same override features for path probabilities that are specified above for branch probabilities.

42.7 Roll Back

The Roll Back Tree Preferences category allows you to control how the model's roll back results are displayed.

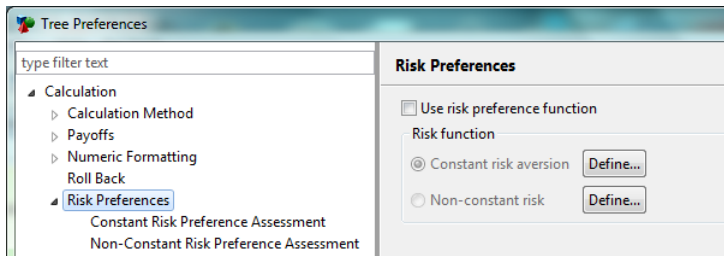


Tree Preferences - Roll Back

- *Display calculated probabilities*: Check this box to display the calculated value of each of the probabilities using numeric formatting preferences during roll back. Turning off roll back will show the original, uncalculated probability expression. This option relates primarily to probabilities which have been entered as variables/expressions.
- *Display EV at terminal and decision nodes & options only*: Check this box to display roll back boxes only at terminal nodes, decision nodes, and branches of a decision node.
- *Fast roll back (no progress bar)*: Check this box to suppress the display of the progress indicator in the status bar. Normally the progress indicator is displayed while a tree is rolling back. Suppressing this display can result in speed increases of up to 100%, depending on the size and complexity of the tree. With either option, you may cancel calculations by pressing ESC.
- *Rollback calculates*: There are four options for the roll back display:
 - Expected values*: Display expected values. This is the default setting.
 - Path Probabilities*: Display path probabilities at all nodes.
 - Payoffs only*: Display payoffs only, rather than expected values at all nodes.
 - Maximin*: Select the most pessimistic option at each chance node and the best option at each decision node.
- *Display path probabilities with all EVs*: Display path probabilities at every node along with the expected values.

42.8 Risk Preferences

The Risk Preferences Tree Preferences category allows you to determine whether to use risk preferences in expected value calculations or not. Refer to the Utility Functions and Risk Preference Chapter for more information.

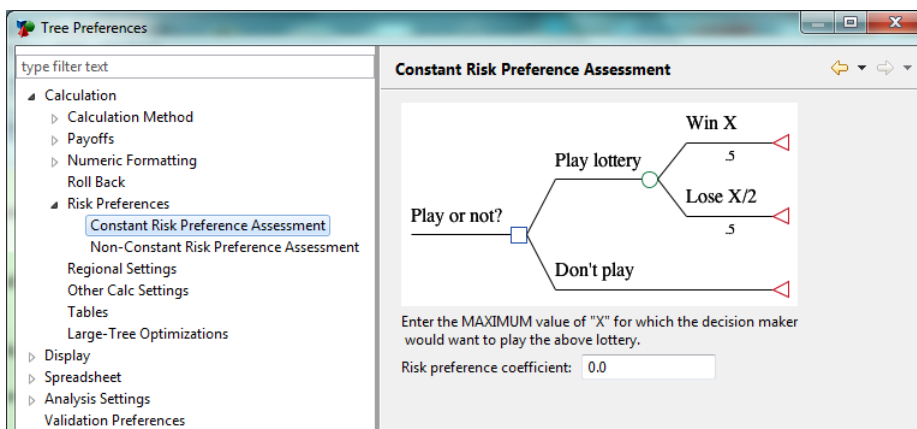


Tree Preferences - Risk Preferences

- *Use risk preference function*: Check this box to calculate the model based on a risk function rather than expected value.
- *Risk function*: Select either a constant or non-constant risk function.

42.8.1 Constant Risk Preference Assessment

The Constant Risk Preference Assessment Tree Preferences category allows you to enter the risk preference coefficient.

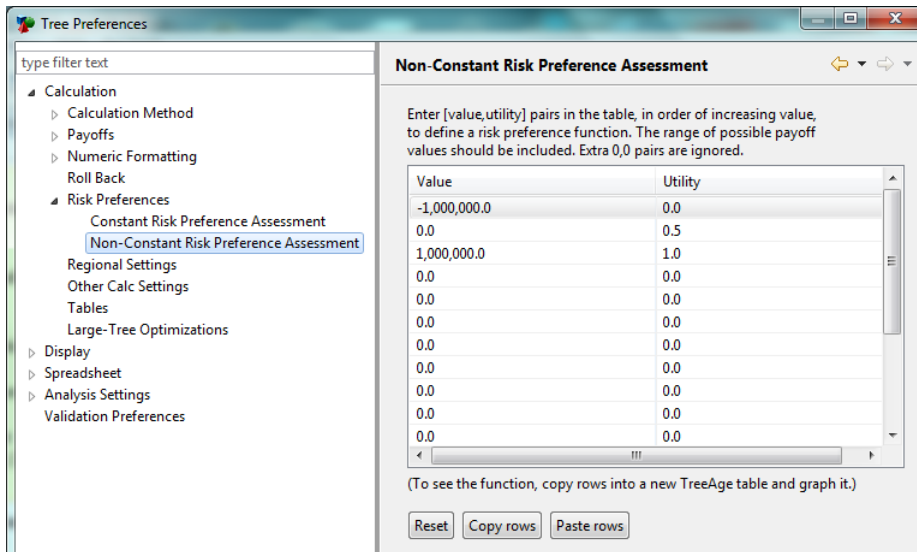


Tree Preferences - Constant Risk Preference Assessment

- *Risk preference coefficient*: Enter a value that defines an amount you are willing to risk for the lottery above. This is used to create a risk preference curve.

42.8.2 Non-Constant Risk Preference Assessment

The Non-Constant Risk Preference Assessment Tree Preferences category allows you to enter a series of risk values to generate a non-constant risk preference curve.



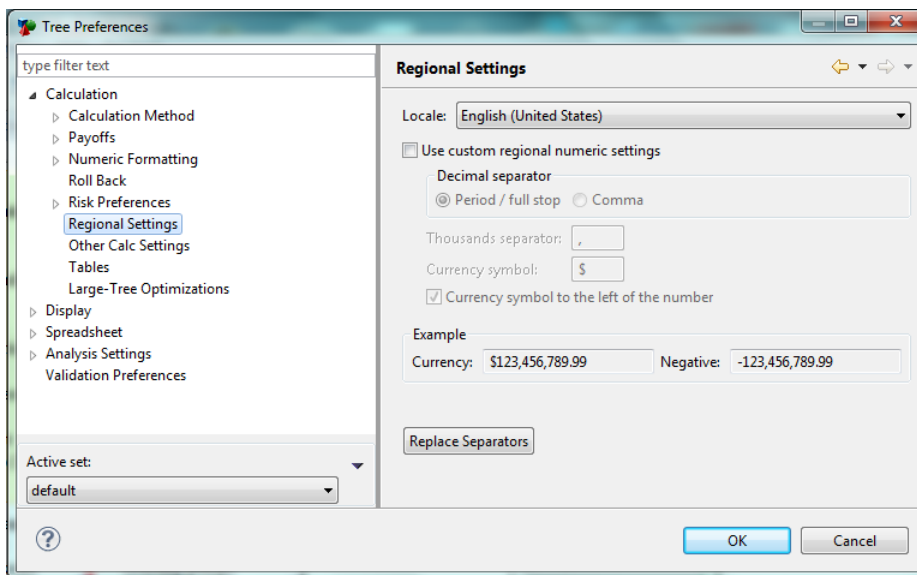
Tree Preferences - Non-Constant Risk Preference Assessment

42.9 Regional Settings

The Regional Settings Tree Preferences category allows you to define regional numeric and currency settings. For new models, these settings are pulled from your computer's operating system. However, you can override the settings in the Tree Preferences; this can be particularly useful when collaborating with colleagues in other countries.



Regional Settings within Tree Preferences control the format of input values (probabilities, payoffs, rewards, variable definitions, etc.). However, the formatting of outputs (roll back, graphs, Monte Carlo output, etc.) is controlled by the computer's regional settings.



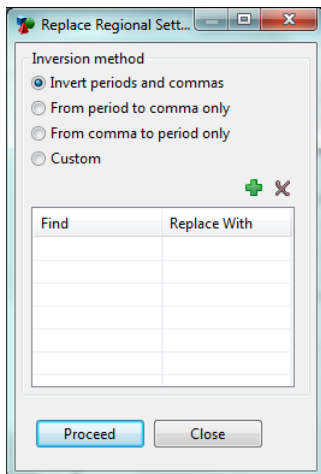
Tree Preferences - Regional Settings

- *Locale*: Select the locale that defines the numeric format for model inputs.
- *Use custom regional numeric settings*: Check this box to override the locale's regional settings for model inputs.
- *Decimal separator*: Select a period (.) or comma (,) as the decimal separator for model inputs.
- *Thousands separator*: Enter a character to use to separate thousands in model inputs. In the US, a comma is used so one million dollars could be entered like this: \$1,000,000.00.
- *Currency symbol*: Enter the currency symbol to use for model inputs.
- *Symbol to the left of the number*: Check this box to enter the currency symbol to the left of the number. Uncheck it to enter the currency symbol to the right of the number.
- *Replace Separators button*: Click this button to replace the separators within the model inputs. This is described in the next section.

42.9.1 Replace Regional Settings

The Replace Regional Settings Tree Preferences category allows you to replace the regional settings within numerical expressions in the model.

These Tree Preferences are enabled only if the "custom regional numeric settings" preference is checked.

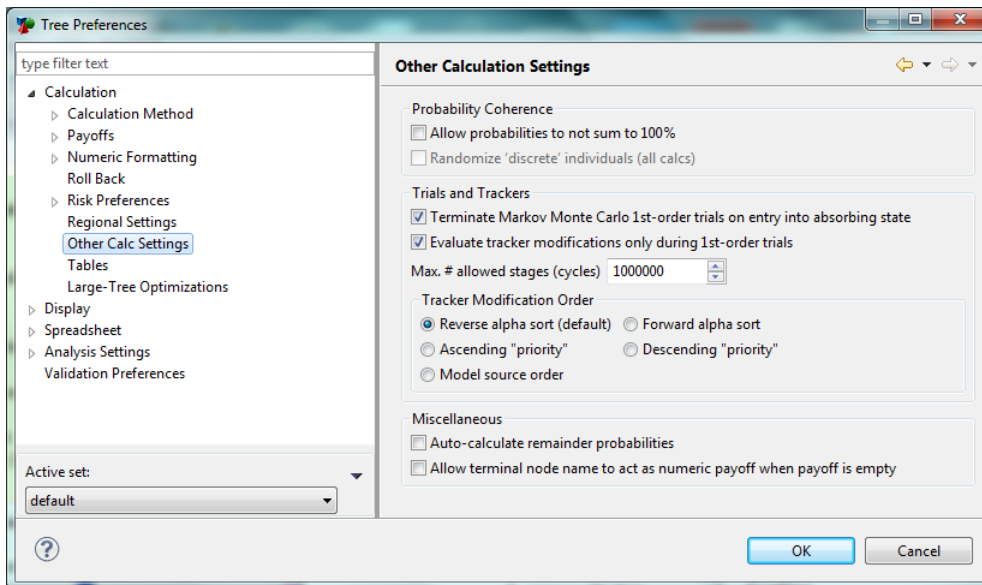


Tree Preferences - Replace Regional Settings

- *Inversion method*: Choose from among the following inversion methods. Each changes the values in numeric expressions within the model.
 - Invert periods and commas*: Replace all periods with commas and vice versa.
 - From period to comma only*: Replace all periods with commas.
 - From comma to period only*: Replace all commas with periods.
 - Custom*: Create custom inversion methods.
- *Find/Replace with grid*: Use the "+" and "X" buttons to add and delete rows from the grid. Enter the target value into the Find column and the replacement value into the Replace With column.

42.10 Other Calc Settings

The Other Calc Settings Tree Preferences category allows you to set a few additional calculation settings.



Tree Preferences - Other Calc Settings

- *Allow probabilities to not sum to 100%:* When checked, TreeAge Pro will not generate an error when probabilities do not sum to 100% and analyses will run. This option should be used only for specific techniques (i.e., dynamic cohort models). Otherwise, this can simply mask errors in the model.
- *Randomize 'discrete' individuals (all calcs):*
- *Terminate Markov Monte Carlo 1st-order trials on entry into absorbing state:* By setting this option, you indicate that the termination conditions should be ignored during Monte Carlo simulations of a Markov process.
- *Evaluate tracker modifications only during 1st-order trials:* When checked, tracker modifications are ignored when not running individual trials through the model. It is only in rare occasions that you would want to uncheck this box.
- *Max. # of allowed stages (cycles):* Set the maximum number of cycles for any Markov model.
- *Tracker Modification Order:* Select the order for tracker modifications to be calculated at a specific node.
- *Auto-calculate remainder probabilities:* When this option is checked, TreeAge Pro will fill in the last probability in a set of branches emanating from a chance node, so long as all the other probabilities on branches emanating from that node are wholly numeric (i.e., no variables are used).
- *Allow terminal node name to act as numeric payoff when payoff is empty:* Check this box to have TreeAge Pro treat the branch description at a terminal node as that node's numeric payoff value.



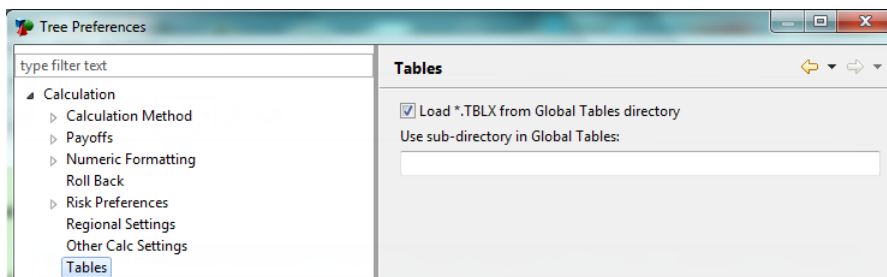
When the Tracker Modification Order sorts by ascending priority, the secondary order is forward alpha. When it is by descending priority, the secondary order is by reverse alpha.

42.11 Large Tree Optimizations

The Large Tree Optimizations tree preferences used in TreeAge Pro 2009 and earlier versions are no longer used starting with TreeAge Pro 2011, due to changes in clone presentation and analysis optimizations.

42.12 Tables

Each tree has its own settings related to the use of Global Tables.



Tables Preferences

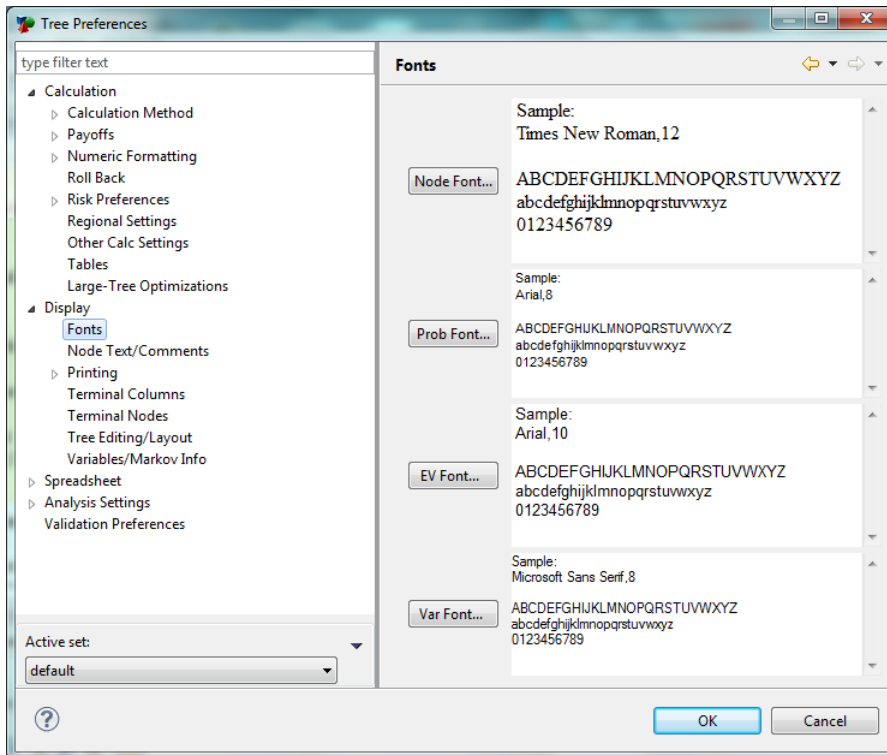
These Tables preferences make it possible to create sub-folders in the Global Tables project specifically for one or for a group of tree models; or, a tree can be set to ignore the Global Tables project and only utilize tables contained within the model.

42.13 Arrows

The Arrows tree display preferences used in TreeAge Pro 2009 and earlier versions is no longer used in later versions of TreeAge Pro. TreeAge Pro now supports individual display settings, including size, for each line/arrow added to the tree.

42.14 Fonts

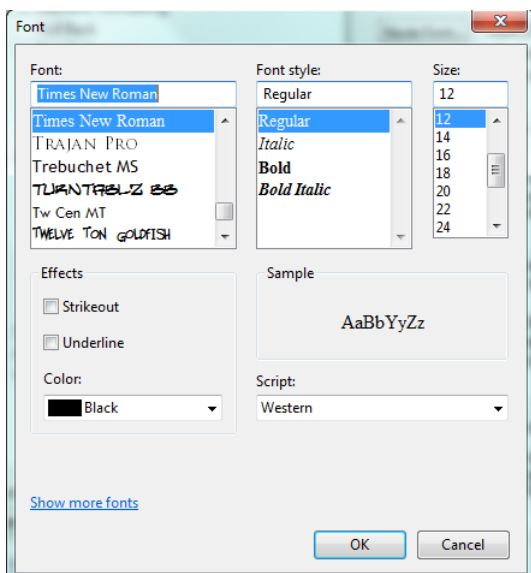
The Fonts Tree Preferences category allows you to specify the font type for several labels/values within the model.



Tree Preferences - Fonts

- *Node Font*: Click this button to select the default font for node labels/text. The font selected in this manner will apply to any new nodes created in the active tree and to any existing nodes, except for any nodes where the font has been set individually.
- *Prob Font*: Click this button to select the default font for probability expressions in both the rolled-back and unrolled-back state. This allows you to clearly distinguish between probability variable names and adjacent node descriptions.
- *EV Font*: Click this button to select the default font for expected value boxes generated during roll back. It also applies to other information which is not user-editable and displayed next to a node, such as clone names when clones are hidden, or payoff names when Always show payoff names is selected.
- *Var Font*: Click this button to select the default font for variable definitions when shown within the tree.

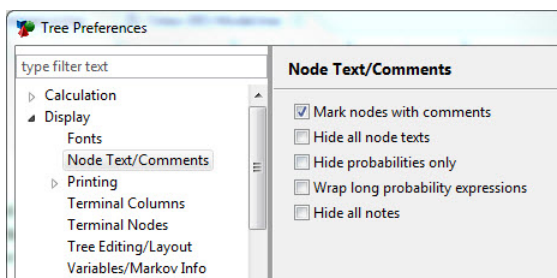
After clicking one of the font buttons, the font selection dialog allows you to specify the font, font style, etc.



Tree Preferences - Fonts - Font Dialog

42.15 Node Text/Comments

The Node Text/Comments Tree Preferences category allows you to show/hide certain text labels/values within the model.

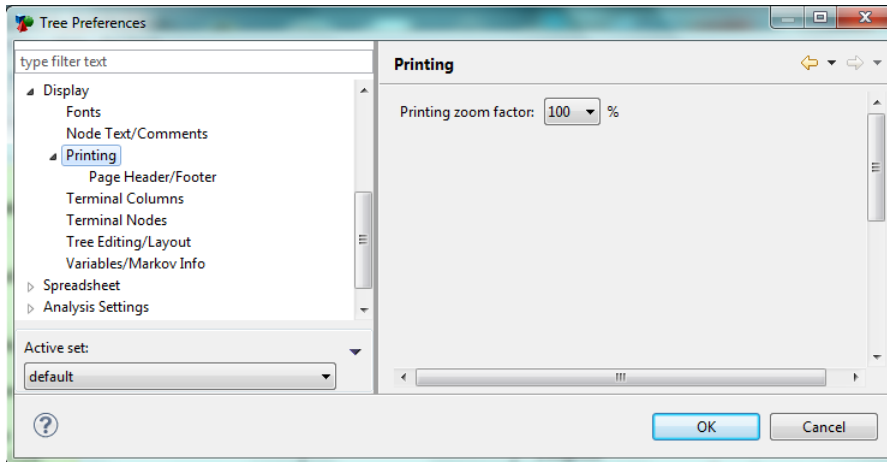


Tree Preferences - Node Text/Comments

- *Mark nodes with comments*: Check this box to display a small flag above the node symbol for all nodes with associated Node Comments. This flag does not print or export.
- *Hide all node texts*: Check this box to suppress all textual information in the Tree Diagram Editor. Use this flag to get a picture of the structure of your tree.
- *Hide probabilities only*: Check this box to hide only probabilities expressions in the Tree Diagram Editor, while all other textual information is visible. Use this flag to temporarily simplify the display of complex trees with many uncertainties.
- *Wrap long probability expressions*: Check this box to wrap probability expressions that may exceed the width of a node.
- *Hide all notes*: Check this box to hide/show all the notes in the Tree Diagram Editor.

42.16 Printing

The Printing Tree Preferences category allows you to specify printing options for the model.



Tree Preferences - Printing

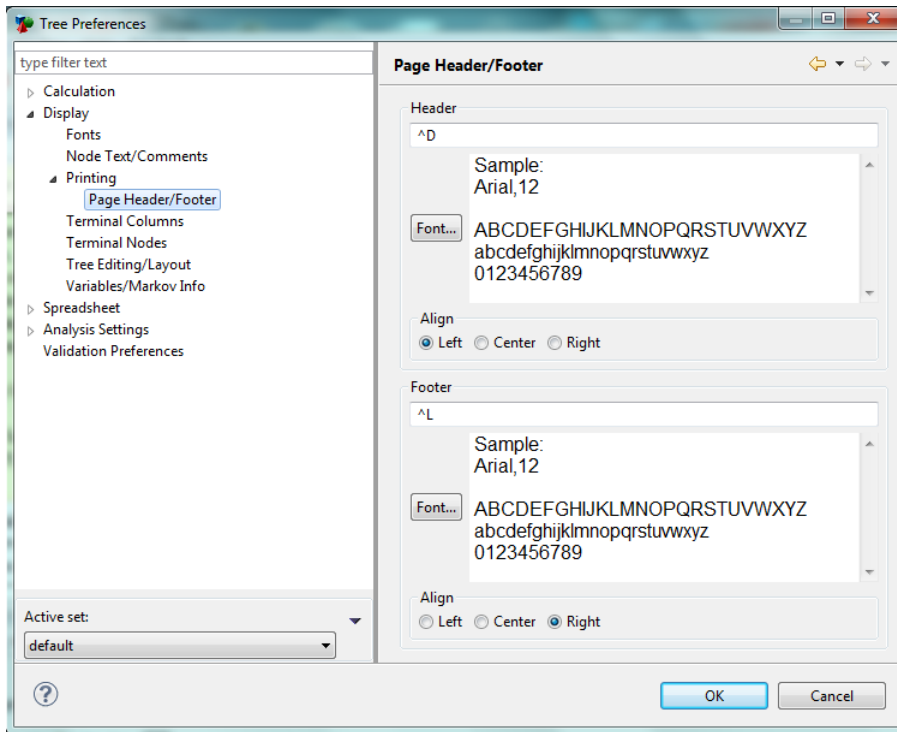
- *Printing zoom factor*. Set the zooming scale for printed output of the model. A lower percentage will fit more of the model on a page.

Note also that it is independent of any scaling specified under Page Setup. Thus, if your printer driver allows scaling via the Page Setup command, you run the risk of applying one percentage against another.

The printing zoom factor is also independent of the screen-display zoom factor, set via the Modeling Palette.

42.16.1 Page Header/Footer

The Page Header/Footer Tree Preferences category allows you to specify a page header and a page footer that are used when printing the model.



Tree Preferences - Page Header/Footer

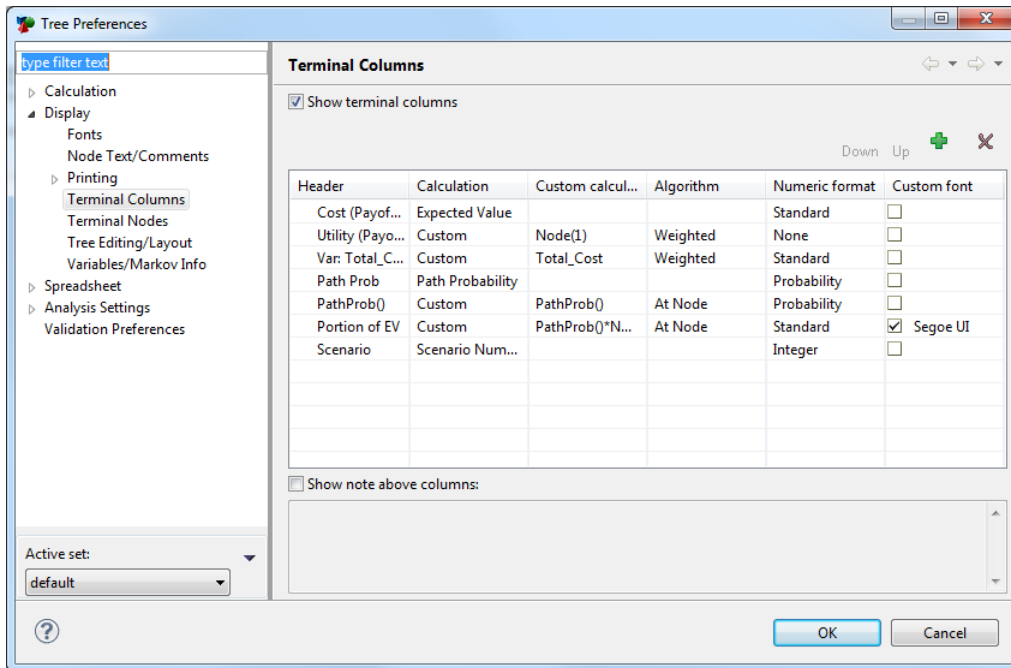
- *Header*: Text to display at the top of each printed page of model.
- *Header Font*: Font to use for header text.
- *Header Align*: Horizontal alignment for header.
- *Footer*: Text to display at the bottom of each printed page of model.
- *Footer Font*: Font to use for footer text.
- *Footer Align*: Horizontal alignment for footer.

42.17 Terminal Columns

The Terminal Columns Tree Preferences category allows you to create terminal columns to display during roll back of the model.

When enabled, terminal columns are presented to the right of the standard payoff values at each terminal node.

The following figure shows the preferences for the the Special Features tutorial example model Terminal Columns.trex.



Tree Preferences - Terminal Columns

- *Show terminal columns*: Check this box to show terminal columns on roll back instead of expected values.
- *Terminal column grid*: Grid showing existing terminal columns.

Terminal columns can be edited directly in the grid. Click the "plus" icon to add a terminal column to the grid. Click the "X" icon to delete the selected terminal column. There is no separate terminal column dialog. The columns in the grid are described below.

- *Header*: Enter text to be displayed above the terminal column data.
- *Calculation*: Select the type of calculation to be performed for the terminal column.
 - *Expected Value*: Show the expected value of the active payoff.
 - *Incremental Value*: Show the incremental change in expected value between this strategy and the previous one.
 - *Path Probability*: Show the path probability for the terminal node.
 - *Scenario Number*: Show the scenario number which counts from one starting at the top terminal node.
 - *Custom*: Show a custom expression as defined in the Custom calculation column.
- *Custom calculation*: Enter a custom expression using TreeAge Pro variables, functions, etc.
- *Algorithm*: Determines how to calculate the value for a collapsed subtree. This only applies to columns using custom calculations.
 - *Weighted*: Use the weighted average from the terminal nodes contained within the collapsed subtree.

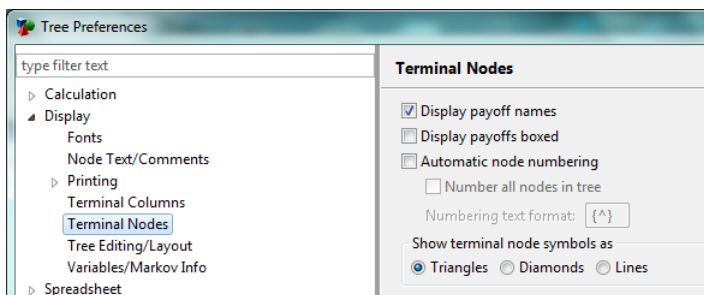
- *At Node*: Use the value from the node at the base of the collapsed subtree.
- *Numeric format*: Select the numeric formatting to use for this value.
 - *Standard*: Use formatting for active payoff.
 - *Probability*: Use formatting for probabilities.
 - *Custom*: Use custom formatting.
- *Custom font*: Change the font preferences for the column.



Terminal columns are discussed in further detail in the Tree Display Preferences and Options Chapter.

42.18 Terminal Nodes

The Terminal Columns Tree Preferences category allows you to specify how terminal nodes are displayed within the model.



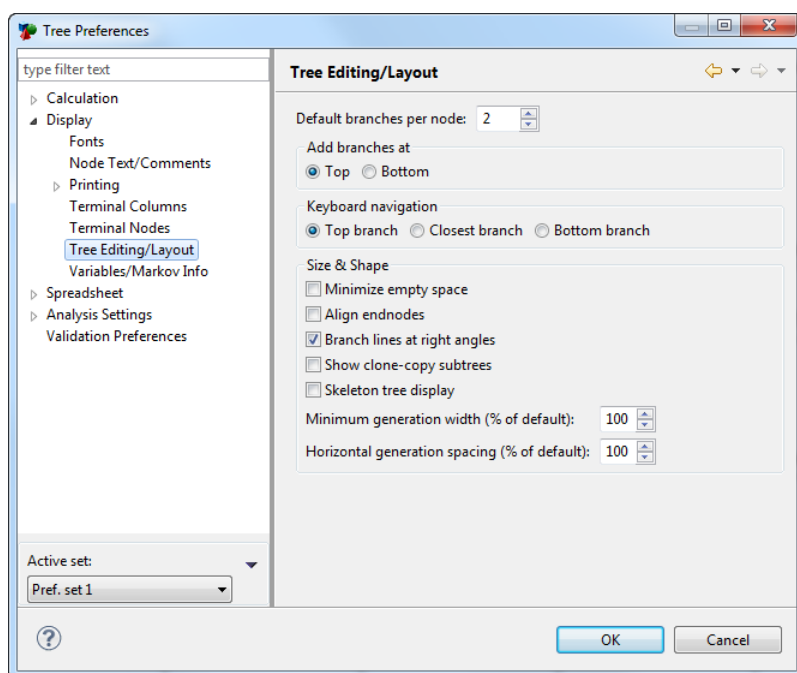
Tree Preferences - Terminal Nodes

- *Display payoff names*: Check this box to display the name/expression of the active payoff at every terminal node. This can be helpful for identifying terminal nodes where a payoff has not been assigned. In the case of trees having multiple payoffs, this feature makes it possible to see at a glance which payoff is active.
- *Display payoffs boxed*: If you have chosen to always display payoff names, check this box to enclose the payoff names/expressions in a box. This option relates only to tree display prior to roll back; during roll back, calculated values are always boxed.
- *Automatic node numbering*: Check this box to display a node number for every terminal node. The numbering format is controlled by the *Numbering text format* field.
- *Number all nodes in tree*: Check this box to number all nodes in the model rather than numbering only the terminal nodes.
- *Numbering text format*: Use this field to specify the custom text for node numbering. Use the ^ (caret) symbol in the text to represent the scenario number.
- *Show terminal node symbols as*: Select one of the three methods for showing terminal nodes in the model. Triangles are the default (and standard) method. Diamonds are used to indicate the

parallelism between terminal nodes in a tree. Lines are for those applications when you do not want any symbol displayed to the right of a final outcome.

42.19 Tree Editing/Layout

The Tree Editing/Layout Tree Preferences category allows you to control a few editing and visual elements of the Tree Diagram Editor.



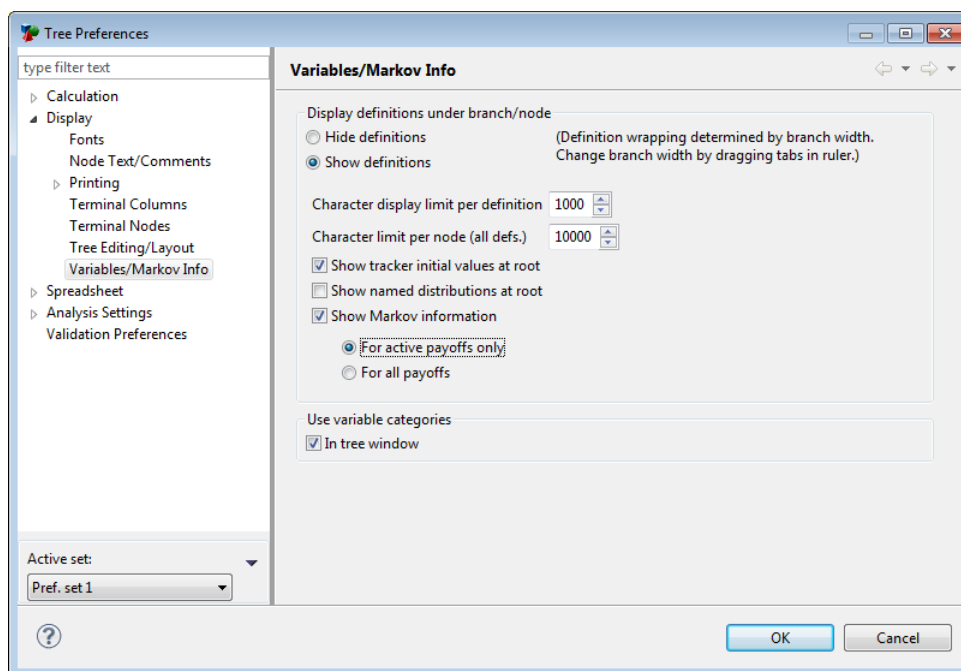
Tree Preferences - Tree Editing/Layout

- *Default branches per node*: Select the number of branches to add when you select Add Branches from the context menu at a node. The default number applies only the first time that branches are added at a given node. Once a node has branches, additional branches are added one at a time.
- *Add branches at*: Select the location for new branches relative to existing branches when adding additional branches to a node. Dragging new nodes from the palette provides additional flexibility in this area.
- *Keyboard navigation*: Select the branch to move to when using the right arrow to navigate from a node to one of its branches.
- *Minimize empty space*: Check this box to produce a “compressed” version of your tree that minimizes empty space. Because each node no longer has its own horizontal “slice” of the tree display, this option may not be used with Align endnodes.
- *Align endnodes*: Check this box to force all terminal nodes to line up at the rightmost edge of the tree.

- *Branch lines at right angles*: Check this box to draw all branch connectors as vertical lines. By default branch connectors are drawn at whatever angle is needed to provide the most direct connection from one node to the next.
- *Show clone-copy subtrees*: Check this box to display clone-copy subtrees; display of clone masters is not affected. When this option is selected, all clone-copy subtrees are displayed with their complete structure. When this option is not selected, only the name of the clone is displayed to the right of the clone-copy anchor node, which can dramatically reduce the visual size of your tree within the Tree Diagram Editor.
- *Skeleton tree display*: Check this box to display and/or print an abstract view of the tree, with space inserted between generations, and collapsed subtrees being completely hidden. Used in combination with the Minimize empty space and Hide clone-copy settings, this creates a highly compact view of the tree.

42.20 Variables/Markov Info

The Variables/Markov Info Tree Preferences category allows you to control the way variables, Markov information and other information is displayed in the Tree Diagram Editor.



Tree Preferences - Variables/Markov Info

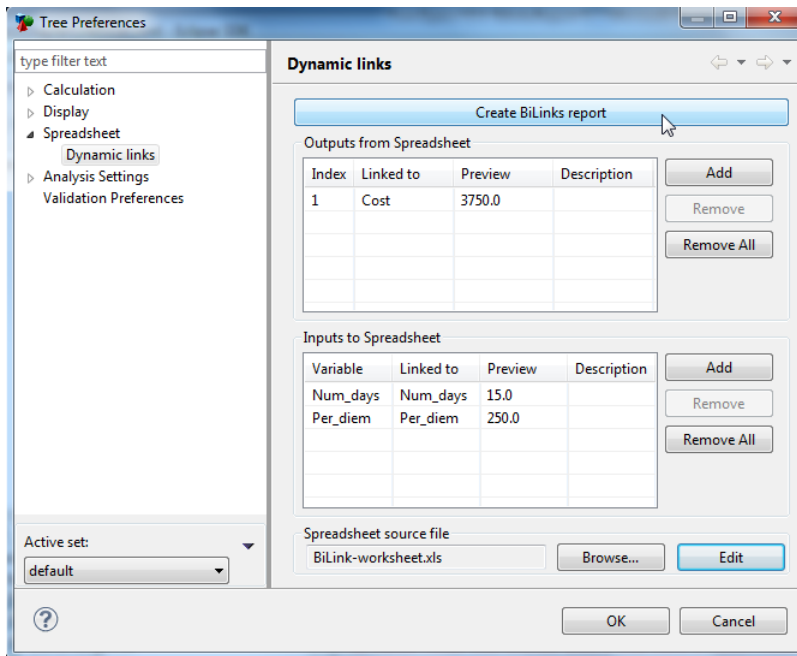
- *Display definitions at nodes*: Select the option to either hide or show variable definitions beneath nodes in the Tree Diagram Editor.
- *Show Markov information*: Show Markov information beneath the node line where applicable (Markov termination condition, Markov state rewards, Markov transition rewards, etc.). Choose whether to display only the active payoffs or all enabled payoffs.
- *Show tracker initial values*: Check this box to show the initial value of trackers at the root node.

- *Show named distributions*: Check this box to show the names of distributions at the root node.
- *Use variable categories in tree window*: Organize variables by categories in the Tree Diagram Editor.

42.21 Dynamic Links

This is the first category related to *spreadsheet* preferences.

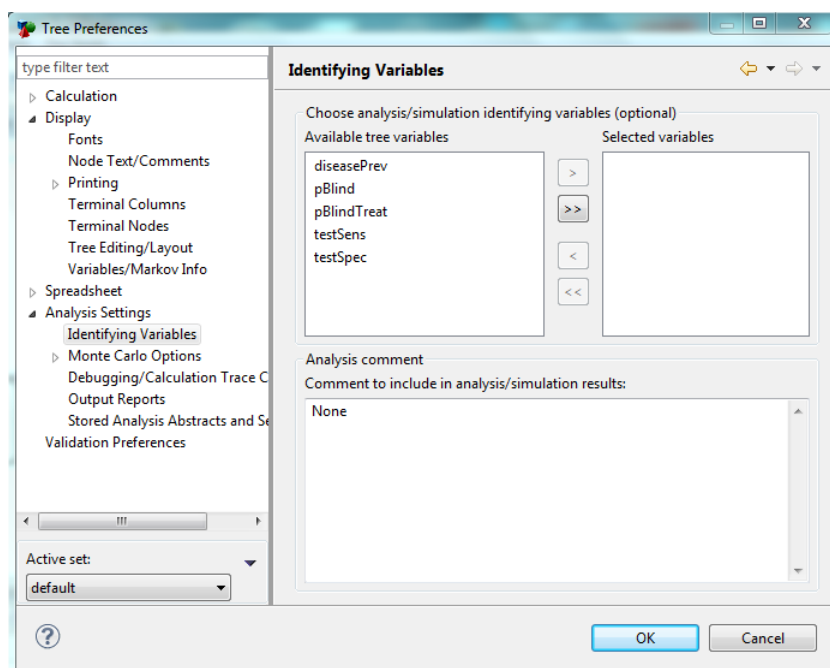
The Dynamic Links Tree Preferences category allows you to edit dynamic links between the model and an Excel workbook. See [Linking with Excel and Other Applications](#) for more details.



Tree Preferences - Dynamic Links

42.22 Identifying Variables

The Identifying Variables Tree Preferences category allows you to store specific variables and their values with Monte Carlo simulation output. This can be useful if you run/save multiple sets of simulation outputs with different variable values - potentially generated by a stored analysis sequence .

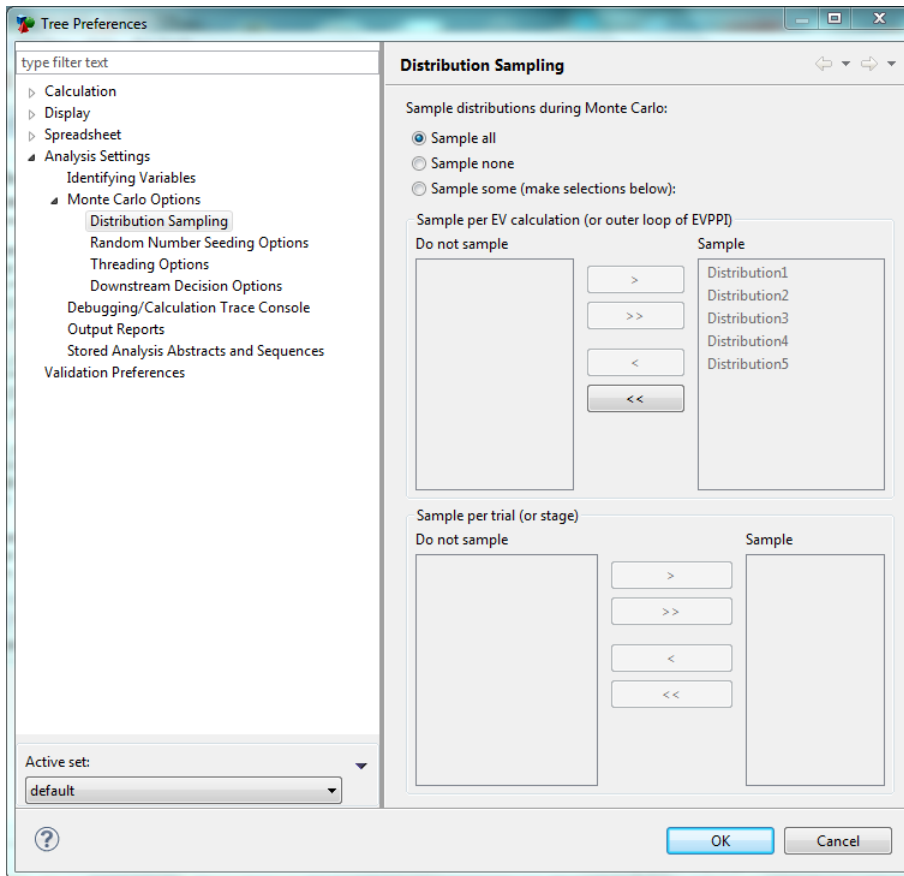


Tree Preferences - Identifying Variables

42.23 Monte Carlo Options

The Advanced Monte Carlo Options Tree Preferences category allows you to specify advanced options for Monte Carlo simulation.

There are four different options found in this category, starting with *Distribution Sampling*.

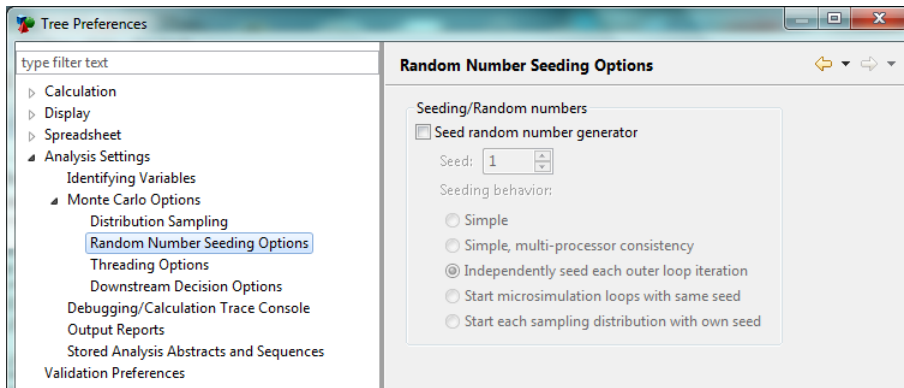


Tree Preferences - Monte Carlo Options - Distribution Sampling

- *Sample all*: Default setting. This will sample all distributions.
- *Sample none*: Check this box so there are no sample distributions.
- *Sample some (make selections below)*: Check this box to choose the distributions you want to sample using the selection boxes.
- *Sample per EV calculation (or outer loop EVPP)*: These selection boxes choose the distributions to sample for the whole cohort.
- *Sample per trial (or stage)*: These selection boxes choose the distributions to sample for each individual trial.

Sample distributions during Monte Carlo:

Random Number Seeding Options

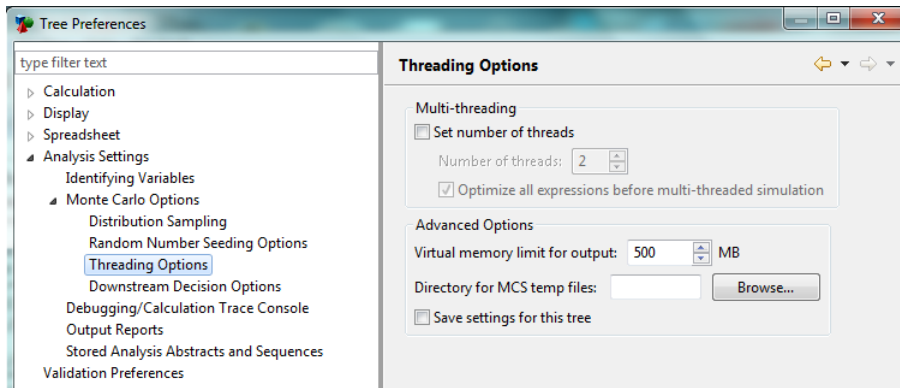


Tree Preferences - Monte Carlo Options - Random Number Seeding Options

- *Seed random number generator*: Check this box to turn seeding on to generate repeatable sampled values from one simulation to the next.
- *Seed*: Choose the initial seed for the random number sequence. Changing this value will generate a different but repeatable series of random numbers.
- *Seeding behavior*: Select which strategy you would like the seed to follow.
- *Simple*: Use a repeatable sequence of numbers.
- *Simple, multi-processor consistency*: This ensures that the sequence of random numbers does not change depending on the number of processors used.
- *Independently seed each out loop iteration*: Default setting. This generates a sequence that does not change if strategies are reordered, removed, or excluded.
- *Start microsimulation loops with same seed*: Uses consistent seeds for inner loops in a multidimensional simulation. This reduces variance among inner loop iterations.
- *Start each sampling distribution with own seed*: This option provides for better consistency in distribution sampling in EVPPI simulations. Without this setting, rerunning a simulation after moving a single distribution from the outer sampling loop to the inner loop (or vice versa) could cause all other, unmoved distributions to nonetheless return new series of sample values. **ONLY USE FOR EVPPI SIMULATIONS.**

Seeding/Random numbers:

Threading Options

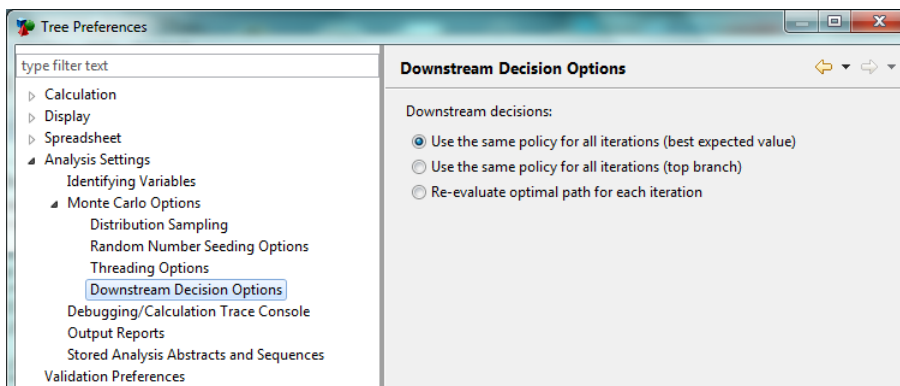


Tree Preferences - Monte Carlo Options - Threading Options

- *Set number of threads*: Check this box to determine a specific amount of threads you wish to run. If unchecked, TreeAge Pro will run a thread on every chip that's available.
- *Virtual memory limit for output*: Set the amount of memory you would like to use. Use this to limit the amount of memory use for the simulation.
- *Directory for MCS temp files*: Choose the location for the temporary files to be written during analysis if necessary.

Multi-threading:

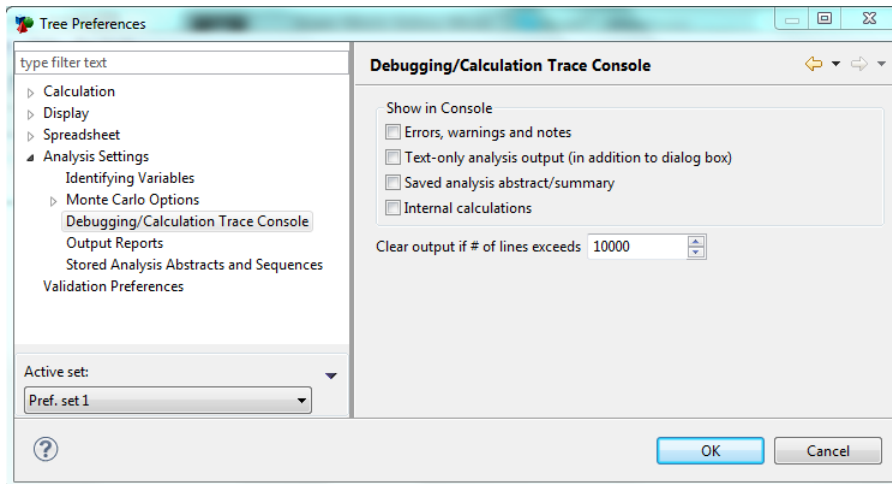
Downstream Decision Options - These options are currently not used.



Tree Preferences - Monte Carlo Options - Downstream Decision Options

42.24 Debugging

The Debugging/Calculation Trace Console Tree Preferences category allows you control over the output in the debug view during analyses.



Tree Preferences - Debugging

- *Show in Console*: Select the items you wish to see in the debug view as TreeAge Pro performs calculations.
- *Errors, warnings and notes*: Check this box to avoid having TreeAge Pro pop-up dialogs when warnings or error messages need to be displayed; instead, checking this box will have the text added to the Calculation Trace Console.
- *Text-only analysis output*: Check this box to write text report analysis output to the Console. For example, Rankings output is purely text and would be written to the Console.
- *Saved analysis abstract/summary*: Check this box to display summary information in the Calculation Trace Console about any new analyses available for storage. This information can be stored for later use via Stored Analysis.
- *Internal calculations*: Check this box to output very detailed, step-by-step results of all tree calculations. This includes variable and function evaluation, payoff and probability calculations, distribution and table references, and more. This is the most commonly useful debugging preference option.
- *Clear output if # of lines exceeds*: Select the maximum number of lines to display within the Calculation Trace Console. Older lines are dropped when the maximum number of lines is exceeded. A larger number holds more outputs in the debug view, which uses more system resources.

42.25 Output Reports

The Output Reports Tree Preferences category allows you to filter the secondary reports available from Monte Carlo simulations.

There are two main options...

- Hide PSA reports for microsimulation
- Display only selected reports from list below

42.25.1 Hide PSA reports for Microsimulation

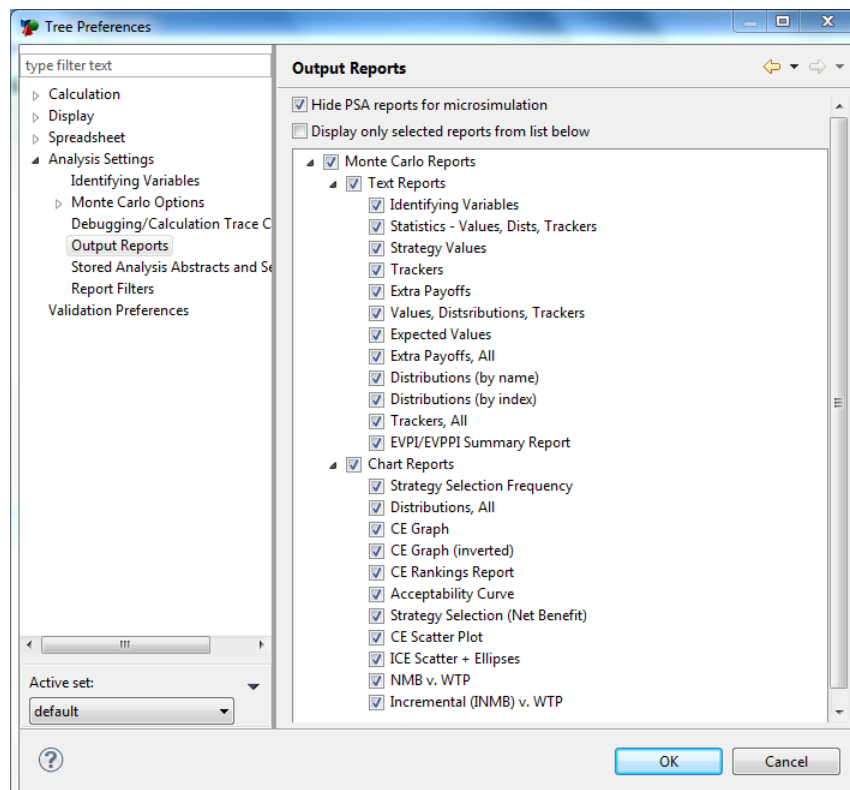
The "Hide PSA reports for microsimulation" option specifically hides all secondary reports that are inappropriate for Microsimulation. Microsimulation is patient-level simulation, so secondary reports are invalid if they rely on incremental calculations among strategies on an iteration-by-iteration basis.

This option defaults to checked. Uncheck the box to see all secondary reports.

42.25.2 Display only selected reports from list below

The "Display only selected reports from list below" option allows modelers to specify which secondary outputs should be displayed. This is normally used when modelers want to simplify the secondary output display to focus on a specific subset of secondary reports.

Check the option "Display only selected reports from list below" to removing some of the output options to make it easier to find the output options you wish to highlight. A modeler might choose to filter the output generated by a Player Model to make it easier for the recipient of the player model to find a specific output.



Tree Preferences - Output Reports

When checked, the "Enable reports filtering" turns on filtering of Monte Carlo simulation output. When unchecked, no filtering is applied.

The group-level checkboxes "Monte Carlo Reports", "Text Reports" and "Chart Reports" options are used to check or uncheck all options within those groups.

The remaining options either include (checked) or exclude (unchecked) that specific output from the Monte Carlo simulation output. Not all options apply to all simulation output. Those restrictions are highlighted below.

- *Identifying Variables* - all output
- *Statistics* - Values, Dists, Trackers - all output
- *Strategy Values* - all output
- *Trackers* - only if model contains trackers
- *Extra Payoffs, All* - only if extra payoffs are included in model
- *Distributions (by name)* - only if model contains distributions
- *Distributions (by index)* - only if model contains distributions
- *Trackers, All* - only if model contains trackers
- *EVPI/EVPPI Summary Report* - all output
- *Strategy Selection Frequency* - only if non-CE model
- *Distributions, All* - only if model contains distributions
- *CE Graph* - CE models only
- *CE Graph (inverted)* - CE models only
- *CE Rankings Report* - CE models only
- *Acceptability Curve* - CE models only
- *Strategy Selection (Net Benefit)* - CE models only
- *CE Scatter Plot* - CE models only
- *ICE Scatter + Ellipses* - CE models only
- *NMB v. WTP* - CE models only
- *Incremental (INMB) v. WTP* - CE models only

Output Report Options

42.26 Stored Analysis Abstracts and Sequences

Stored Analyses and Sequences are described in a separate chapter.

42.27 Application Preferences

Application Preferences control many settings and options for the TreeAge Pro application and are not associated with a specific model, and therefore apply to all models.

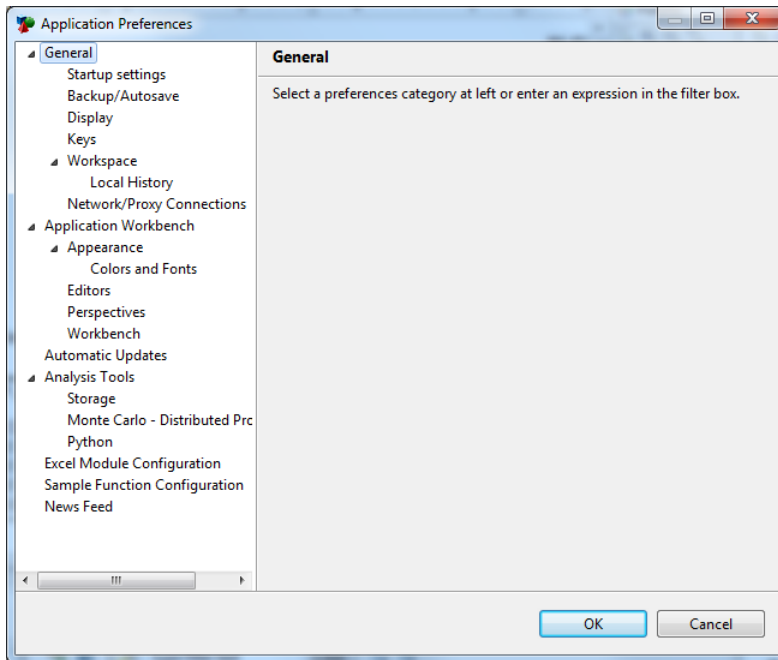
Application Preferences are edited using the Application Preferences Dialog.

42.28 Application Preferences Dialog

The Application Preferences Dialog is used to edit Application Preferences.

To open the Application Preferences Dialog:

- Choose Window > Application Preferences from the menu.

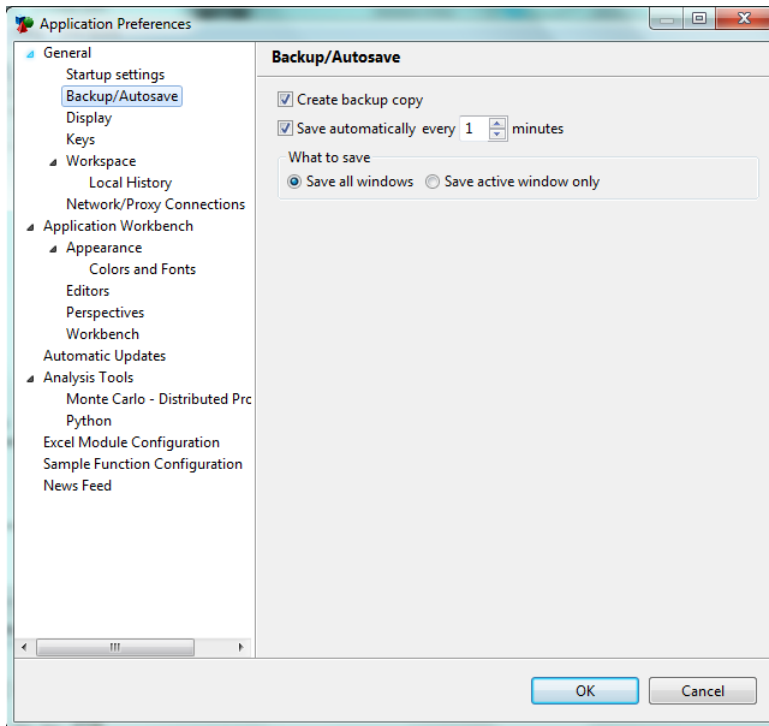


Application Preferences Dialog

Subsequent sections will describe each category of Application Preferences.

42.29 Application Preferences - Backup/Autosave

The Backup/Autosave Application Preferences category allows you control on how TreeAge Pro backs up your work.



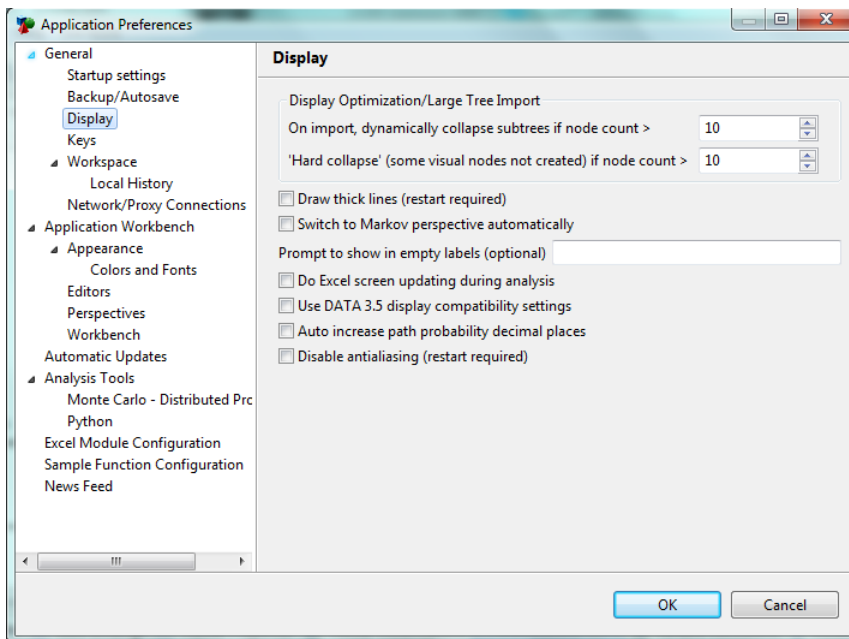
Application Preferences - Backup/Autosave

- *Create backup copy*: Check this box, and TreeAge Pro will save a backup copy of each file that you open and subsequently save changes to.
The backup file contains the version of the file as it existed when it was opened. No matter how many times you modify or save the file after opening it, its backup file continues to contain the original version of the document. If you close a file and then reopen it, the backup process is restarted; if a backup file already exists for a document that you open, TreeAge Pro will overwrite the existing backup file when you save the document.
The backup file is created in the same directory as the original file, with “-backup” appended to the extension; for example, the backup file for “my tree #1.tre” would be saved as “.my tree #1.tre-backup”.
If you accidentally save changes to a file that cannot be undone, you can open the backup copy of the file. To do so, however, you will need to rename the backup file as a *.trex file first. Change the Open File dialog’s “Files of type:” drop-down menu to “All files (*.*)”.
- *Save automatically every ____ minutes*: Check this box and TreeAge Pro will automatically save open documents to the original filename at regular intervals without prompting. Only documents that have unsaved changes will be autosaved. Select the number of minutes after which an autosave file should be generated.

- *What to save*: Select the option to create autosave files for all open models or only for the active model.

42.30 Application Preferences - Display

The Display Application Preferences category allows you to control how the application displays all models.

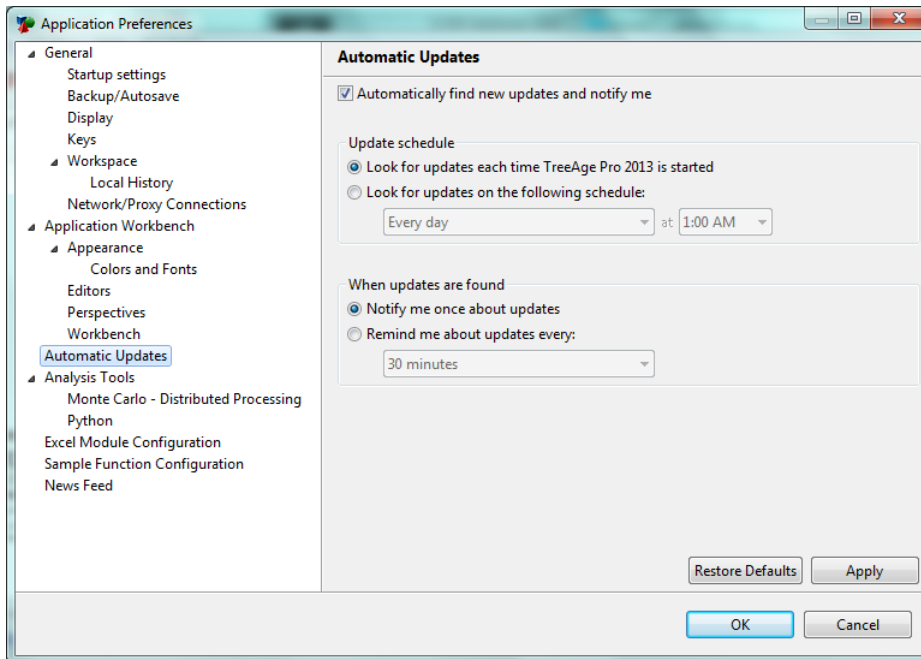


Application Preferences - Display

- *Display Optimization/Large Tree Import*:
- *On import, dynamically collapse subtrees if node count > __*:
- *'Hard collapse' (some visual nodes not created) if node count > __*:
- *Draw thick lines (restart required)*:
- *Switch to Markov perspective automatically*:
- *Prompt to show in empty labels (optional)*:
- *Do Excel screen updating during analysis*:
- *Use DATA 3.5 display compatibility settings*:
- *Auto increase path probability decimal places*:
- *Disable antialiasing (restart required)*:

42.31 Application Preferences - Automatic Updates

The Automatic Updates Application Preferences category allows you control over how the TreeAge Pro application is updated as new updates are released.

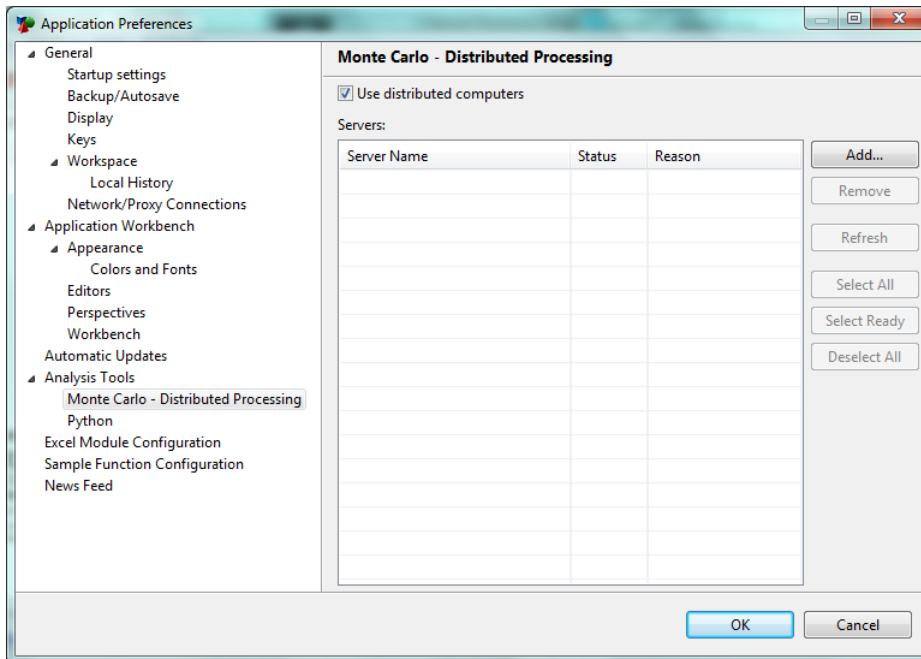


Application Preferences - Automatic Updates

- *Automatically find new updates and notify me*: Check this box to have TreeAge Pro automatically check for new updates to the software as they become available.
- *Update schedule*: Select an option for when to check for new updates.
Look for updates each time TreeAge Pro is started: Check each time TreeAge Pro is started.
Look for updates on the following schedule: Check based on a custom schedule.
- *When updates are found*: Select a schedule for notifications regarding available updates.
Notify me once about updates: Notify once when updates are available but never again.
Remind me about updates every: Notify repeatedly when updates are available based on a custom schedule.

42.32 Application Preferences - Monte Carlo Distributed Processing

The Monte Carlo Distributed Processing Application Preferences category allows you to setup helper computers to speed up Monte Carlo simulations.

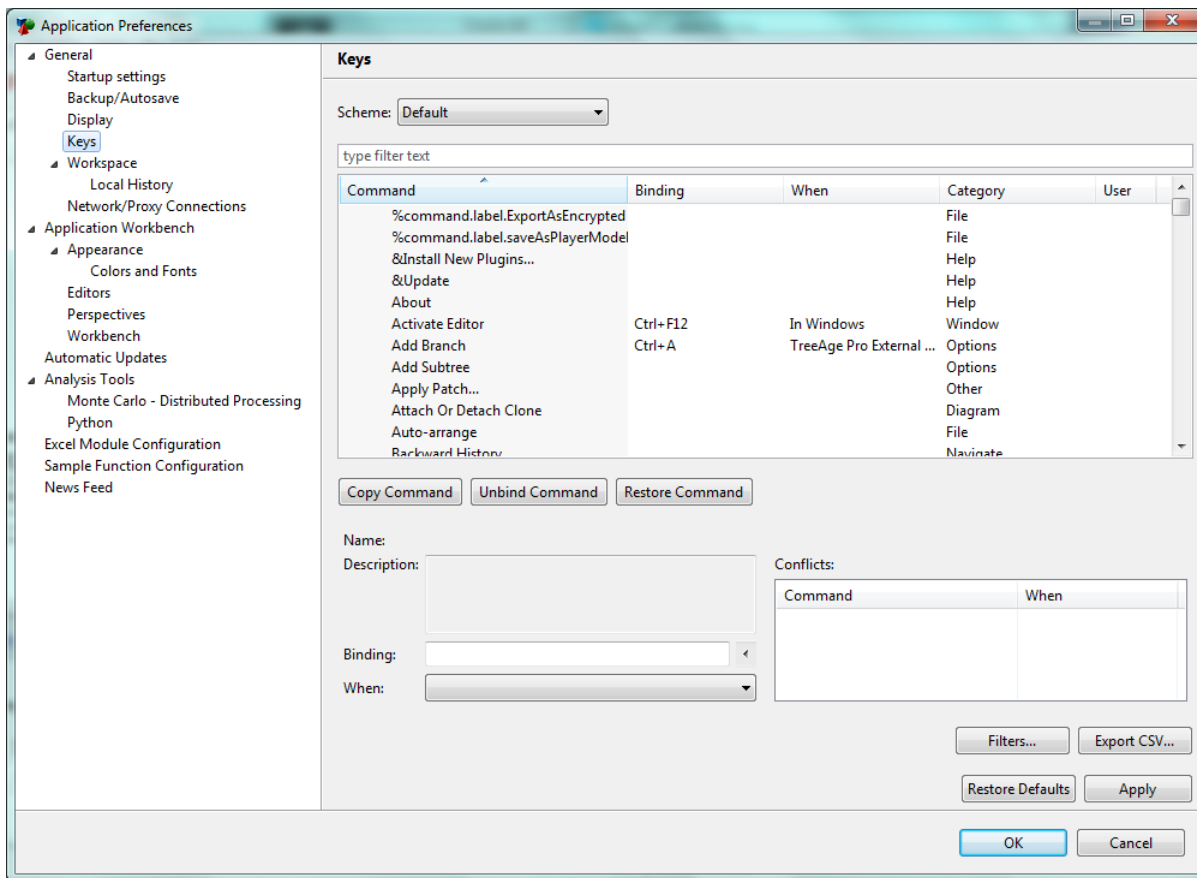


Application Preferences - Keys

- *Use distributed computers*: Check this box to use helper computers.
- *Server name*: The helper computer name. The helper computer can be used to perform some iterations of a Monte Carlo simulation so that the overall simulation runs faster.
- *Status*: The status of the helper computer, which indicates whether the helper computer is available.
- *Reason*: Purpose for which the helper computer was added.
- *Add button*: Add a new helper computer.
- *Remove button*: Remove the selected helper computer.
- *Refresh button*: Refresh the status of all helper computers.
- *Select All button*: Select all helper computers in the grid.
- *Select Ready button*: Select all helper computers that are ready.
- *Deselect All button*: All helper computers are no longer selected.

42.33 Application Preferences - Keys

The Keys Application Preferences category allows you to setup keyboard shortcuts for functions within TreeAge Pro.

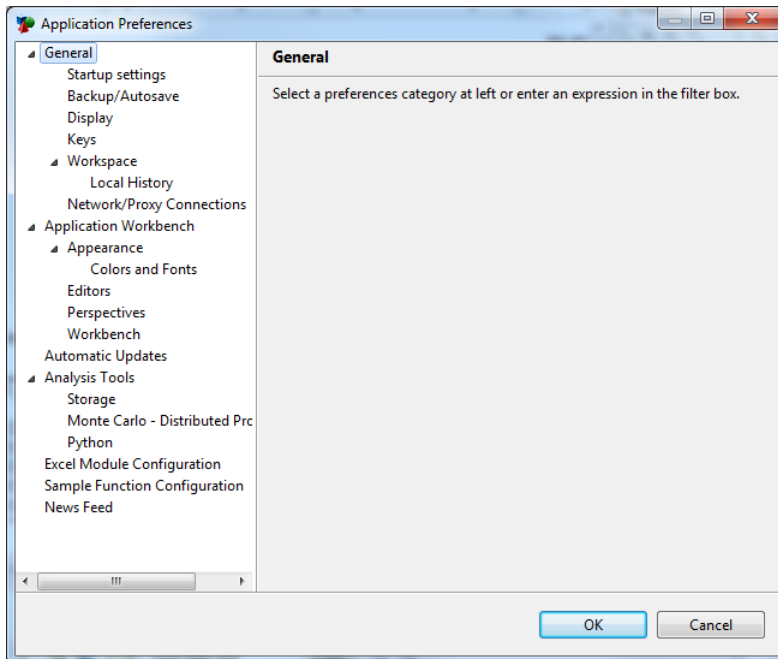


Application Preferences - Keys

-

42.34 Application Preferences - Storage

The Storage Application Preference category allows you to store analysis data on disk rather than in memory. Note that this option is only available for 64-bit versions of TreeAge Pro.



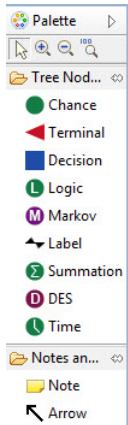
Application Preferences - Storage

43. Technical Details & Utilities

This chapter contains technical details on the TreeAge Pro application and TreeAge Pro utilities.

43.1 Node Types

Decision trees can include nodes with a number of different node types. This section lists the node types and their typical usage. The Tree Diagram Editor Palette displays all the available node types.



Tree Diagram Editor Palette

- *Chance*: Used to represent a situation with an unknown outcome. Each branch of the chance node represents one of the outcomes. There must be a branch for every possible outcome, and each branch will include a probability expression for the likelihood of that outcome. The hashtag (#) can be used for one of the branch probabilities, which is calculated as the complement of the sum of all other probability expressions.
- *Terminal*: Used to terminate endpoints in a model. All endnodes (no branches) must be terminal nodes. In a decision tree, a terminal node represents the end of a scenario, and payoff values are required to place value(s) on that scenario. In a Markov model, a terminal node represents the end of a cycle, and a jump state is used to return the cohort to a health state to start the next cycle. In DES models, terminal nodes function the same as in Markov models.
- *Decision*: Used to represent a decision point within a model. Each branch represents a strategy which will be evaluated separately to determine the optimal strategy at that decision node. It is typical for a model to contain a single decision node as the left most node in the model (the root node).
- *Logic*: Used like a chance node with an unknown outcome. However, each branch will have a logical expression instead of a probability expression. Logical expressions are checked from the top branch down. The model will flow to the first branch with a logical expression that evaluates to true. It is common to use the hashtag (#) for the bottom branch for when all logical expressions for the other branches are false.

- *Markov*: Used as the start of a Markov model. The Markov node and its entire subtree are part of the Markov model, which is normally part of a larger decision tree model.
- *Label*: Used to simply flow from a node to its one and only branch. It functions the same as a chance node with a single branch with probability of 100%. This is commonly used to separate dependent calculations, so the calculations at the label node are complete before the calculations are performed at the branch node.
- *Summation*: Used to sum the expected values (EVs) of all branches within a decision tree, rather than use a weighted average as is done for chance nodes. This is commonly used when there are multiple separate value calculations and the sum of those values represents the total EV. For example, branches could be used in a legal model for compensatory damages and punitive damages, such that the total value is the sum.
- *DES*: Used as the start of a Discrete Event Simulation (DES) model. The DES node and its entire subtree are part of the DES model, which is normally part of a larger decision tree model.
- *Time*: Used only within a DES model. Each branch of a DES node represents a possible event with a time-to-event expression. Time passes and time-based value is accumulated at this node until an event occurs.

Node Types

43.2 Random number generator details

Monte Carlo simulations in TreeAge Pro make use of a robust pseudo-random number generator (RNG) algorithm, the Mersenne Twister, which has the following useful properties:

1. Has a period of 2^{19937} , or approximately 10^{6001} unique sequences.
2. Has negligible serial correlation between successive values in the output sequence.
3. Works fast.
4. Passes numerous tests for statistical randomness.

Distribution sampling and discrete simulation random walks utilize the RNG. By default, each RNG is “seeded” using the computer clock. This “random” seeding can be overridden by the user specifying a random seed value in the Simulation options.

In a multi-processor simulation, each thread has a separate RNG, which is started at a different position based on the clock seed or the user-specified.

43.3 Report Output Files (*.rptx)

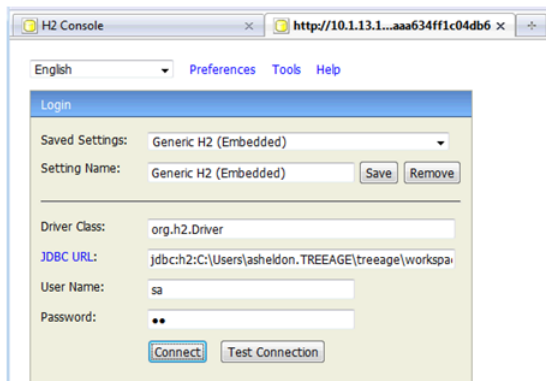
Report outputs from Monte Carlo, Sensitivity, and Markov Cohort analyses can be saved in *.rptx files. These files can be reopened in TreeAge Pro to generate graphical and text output.

The *.rptx file is a zip file, which contains an H2 datasource/database (plus other files). This datasource is queryable from outside of TreeAge Pro through an H2 browser console.

You can copy the *.rptx files, change the extension to *.zip, and uncompress the files ("datasource" and "datasource.*" files are needed). The path to "datasource" will be used to connect in the browser console.

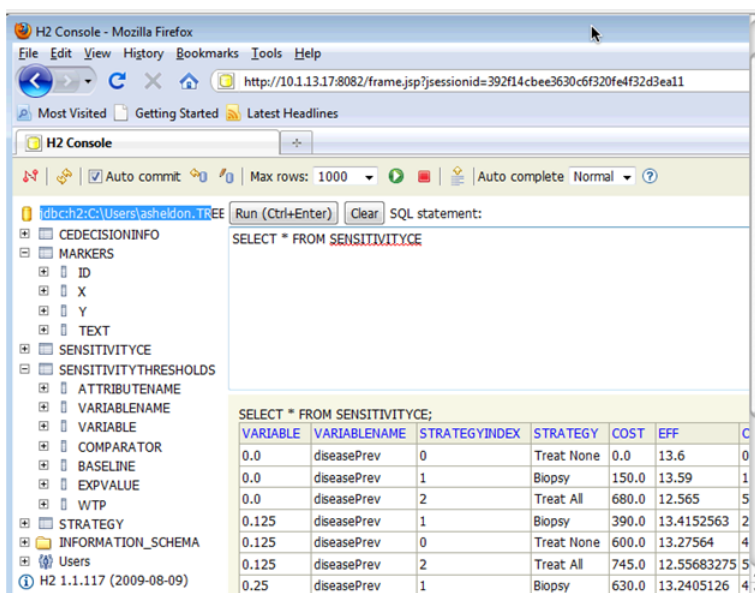
It is possible to then run the H2 database service/console from an "h2.jar" file zipped within the TreeAge Pro plugins directory; it will be easier to install H2 separately. Click here to go to the H2 site. The Download page will allow you to download the H2 Database Engine. The Quickstart page provides information on installation.

You can then run the h2.jar to open the console in a browser, add/browse to the path of your datasource, and enter the username ("sa") and password ("sa").



H2 Login within browser

The H2 Console then provides access to the database tables, allowing you to create queries to pull data as needed.



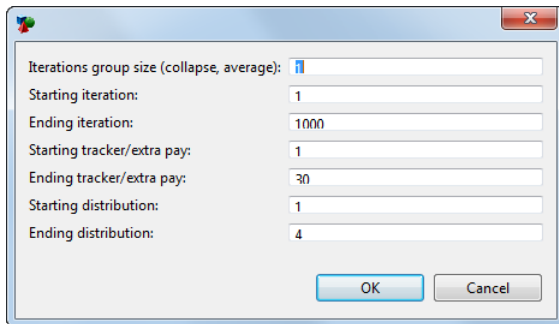
H2 Console within browser

43.4 Importing TP 2009 Simulation Output

The current TreeAge Pro software can import the *.MCS simulation output files generated by TreeAge Pro 2009 and earlier software versions.

To import an *.MCS file:

- Choose File > Import/Export > Import TreeAge 200X MCS File... from the menu.
- Enter options in the Import MCS dialog (see below) and click OK.

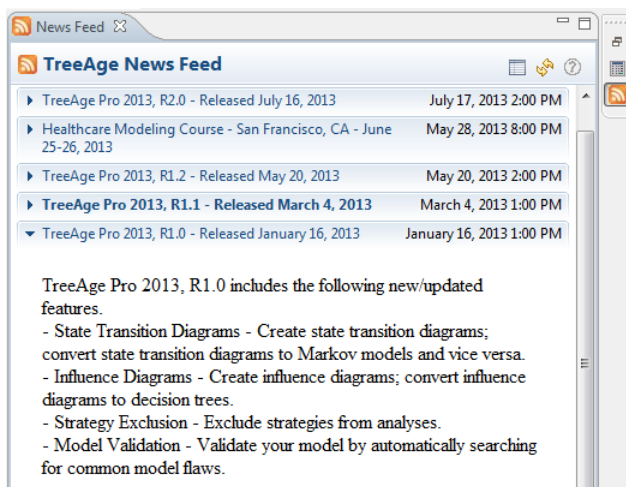


Import MCS Dialog

The options in the dialog above allow you to filter the number of rows and/or columns to include in the imported output. This can reduce the size of the data imported.

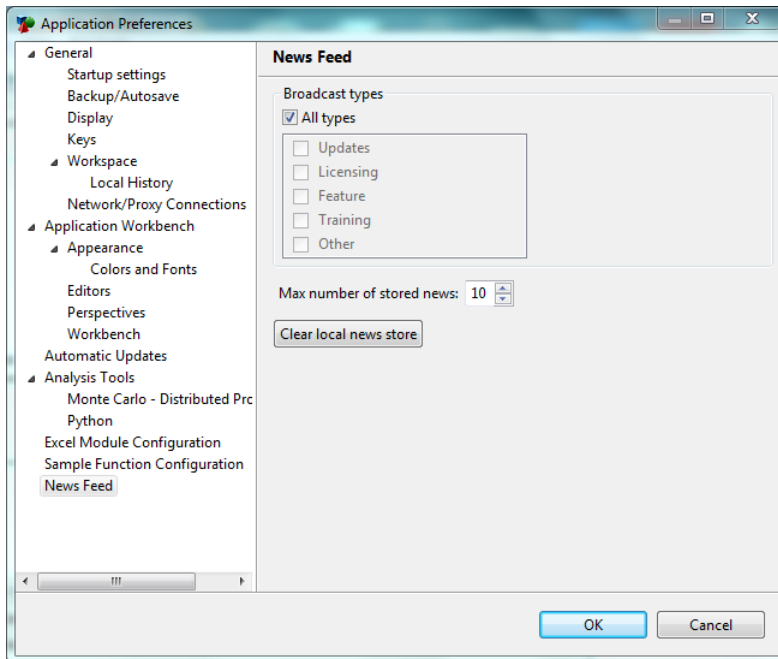
43.5 TreeAge Pro News Reader

The News Reader is used to provide TreeAge Pro users with important information regarding product updates, new products, training, licensing issues, etc. When new news items are available, the News Reader will retrieve and display the items.



News Reader View

Under Window, there are Application Preferences available to control the type of news items you wish to see.



News Feed Application Preferences